IC SEQUENCES AND PROTEINS RELATED TO ALZHEIMER'S DISEASE

RELATED APPLICATIONS

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This application is a continuation-in-part of U.S. Patent Application Serial No. 08/496,841, filed June 28, 1995, which is a continuation-in-part of U.S. Patent Application Secial No. 08/431,048, filed April 28, 1995.

FIELD OF THE INVENTION

The present invention relates generally to the field of neurological and physiological dysfunctions associated with Alzheimer's Disease. More particularly, invention is concerned with the identification, isolation and cloning of the gene which when mutated is associated with Alzheimer's Disease as well as gene products and associated sequence transcript. The present genes. information and neighbouring invention also relates to methods of diagnosing for and detection of carriers of the gene, Alzheimer's Disease diagnosis, gene therapy using recombinant technologies and therapy using the information derived from the DNA, protein, and the metabolic function of the protein. BACKGROUND OF THE INVENTION

In order to facilitate reference to various journal articles, a listing of the articles is provided at the 25 end of this specification.

degenerative is Alzheimer's Disease (AD) central nervous system disorder of the human memory impairment and characterized by progressive cognitive and intellectual decline during mid to late adult life (Katzman, 1986). The disease is accompanied constellation of neuropathologic features by presence amongst which are the principal

or senile plaques and extracellular amlyoid neurofibrillary degeneration of neurons. The complex, although in is disease families it appears to be inherited as an autosomal However, even among these inherited dominant trait. forms of AD, there are at least three different genes which confer inherited susceptibility to this disease (St. George-Hyslop et al., 1990). The 4 (Cys112Arg) allelic polymorphism of the Apolipoprotein E (AopE) gene has been associated with AD in a significant proportion of cases with onset late in life (Saunders et al., 1993; Similarly, a very small Strittmatter et al., 1993). proportion of familial cases with onset before age 65 years have been associated with mutations in the \mathcal{B} amyloid precursor protein (APP) gene (Chartier-Harlin et al., 1991; Goate et al., 1991; Murrell et al., 1991; Karlinsky et al., 1992; Mullan et al., 1992). A third locus (AD3) associated with a larger proportion of cases with early onset AD has recently been mapped to chromosome 14q24.3 (Schellenberg et al., 1992; St. George-Hyslop et al., 1992; Van Broeckhoven et al., 1992).

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Although chromosome 14q carries several genes which could be regarded as candidate genes for the site of mutations associated with AD3 (e.g. cFOS, alpha-1-antichymotrypsin, and cathepsin G), most of these candidate genes have been excluded on the basis of their physical location outside the AD3 region and/or the absence of mutations in their respective open reading frames (Schellenberg, GD et al., 1992; Van Broeckhoven, C et al., 1992; Rogaev et al., 1993; Wong et al., 1993).

There have been several developments and commercial treatment of Alzheimer's respect of in directions Published diagnosis thereof. disease and application WO 94 23049 describes transfection of high molecular weight YAC DNA into specific mouse cells. This method is used to analyze large gene complexes, for example the transgenic mice may have increased amyloid precursor protein gene dosage, which mimics the trisomic condition that prevails in Downs Syndrome and the generation of animal models with &-amyloidosis prevalent individuals with Alzheimer's Disease. Published describes 00569 94 application international WO transgenic non-human animals harbouring large trans genes such as the trans gene comprising a human amyloid precursor protein gene. Such animal models can provide useful models of human genetic diseases such as Alzheimer's Disease.

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Canadian Patent application 2096911 describes a nucleic acid coding for amyloid precursor protein-cleaving protease, which is associated with Alzheimer's Disease and Down's syndrome. The genetic information may be used to diagnose Alzheimer's disease. The genetic information was isolated from chromosome 19. Canadian patent application 2071105, describes detection and treatment of inherited or acquired Alzheimer's disease by the use of YAC nucleotide sequences. The YACs are identified by the numbers 23CB10, 28CA12 and 26FF3.

U.S. Patent 5297562, describes detection of 30 Alzheimer's Disease having two or more copies of chromosome 21. Treatment involves methods for reducing the proliferation of chromosome 21 trisomy. Canadian Patent Application 2054302, describes monoclonal antibodies which recognize human brain cell nucleus protein encoded by chromosome 21 and are used to detect changes or expression due to Alzheimer's Disease or Down's Syndrome. The monoclonal antibody is specific to a protein encoded by human chromosome 21 and is linked to large pyramidal cells of human brain tissue.

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extensive effort and a unique approach investigating the AD3 region of chromosome the gene has Alzheimer's related membrane protein (ARMP) been isolated, cloned and sequenced from within the AD3 In addition, direct region on chromosome 14g24.3. sequencing of RT-PCR products spanning this 3.0 kb cDNA transcript isolated from affected members of at least eight large pedigrees linked to chromosome 14, has led to the discovery of missence mutations in each of these different pedigrees. These mutations are absent in normal chromosomes. It has not been established that the ARMP gene is causative of familial Alzheimer's In realizing this link, Disease type AD3. understood that mutations in this gene can be associated with other cognitive, intellectual, or psychological such as cerebral hemorrhage, schizophrenia, disease depression, mental retardation and epilepsy. phenotypes are present in these AD families and these phenotypes have been seen in mutations of the APP protein gene. The Amyloid Precursor Protein (APP) gene is also associated with inherited Alzheimer's Disease. The identification of both normal and mutant forms of the ARMP gene and gene products has allowed for the development of screening and diagnostic tests for ARMP utilizing nucleic acid probes and antibodies to the gene product. Through interaction with the defective gene product and the pathway in which this gene product is involved, gene therapy, manipulation and delivery are now made possible.

SUMMARY OF THE INVENTION

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Various aspects of the invention are summarized as In accordance with a first aspect of the follows. is mammalian polynucleotide purified a invention, provided which codes for Alzheimer's related membrane protein (ARMP). The polynucleotide has a sequence which is the functional equivalent of the DNA sequence of ATCC deposit 97124, deposited April 28, 1995. The mammalian polynucleotide may be in the form of DNA, genomic DNA, CDNA, mRNA and various fragments and portions of the gene sequence encoding ARMP. The mammalian DNA is conserved in many species, including human and rodents, example, mice. The mouse sequence encoding ARMP has greater than 95% homology with the human sequence encoding the same protein.

Purified human nucleotide sequences which encode mutant ARMP have mutations at nucleotide position i) 685, A→C ii) 737, A→G iii) 986, C→A, iv) 1105, C→G, v) 1478, G→A, vi) 1027, C→T, vii) 1102, C→T and viii) 1422, C→G of Sequence ID No: 1 as well as in the cDNA sequence of a further human clone of a sequence identified by ID NO:133.

The nucleotide sequences encoding ARMP have an alternative splice form in the genes open reading frame.

The human cDNA sequence which codes for ARMP has

sequence ID No. 1 as well as sequence SEQ ID NO:133 as sequenced in another human clone. The mouse sequence which encodes ARMP has SEQ ID NO:3, as well as SEQ ID NO:135 derived from a further clone containing the entire coding region. Various DNA and RNA probes and primers may be made from appropriate polynucleotide lengths selected from the sequences. Portions of the sequence also encode antigenic determinants of the ARMP.

Suitable expression vectors comprising the nucleotide sequences are provided along with suitable host cells transfected with such expression vectors.

In accordance with another aspect of the invention, purified mammalian Alzheimer's related membrane protein The purified protein has an amino acid is provided sequence encoded by polynucleotide sequence 15 identified abave which for the human is SEQ ID NO:2 and SEQ ID NO:134 $\sqrt{\text{derived from another clone}}$. The mouse amino acid sequence is defined by SEQ ID NO:2 and SEQ ID NO.136, the late \frak{t} being translated from another clone containing the entire coding region. The purified 20 protein may have Aubstitution mutations selected from the group consisting of positions identified in SEQ ID NO:2 and Sequence ID NO:134.

- i) M 146L
- 25 ii) H 163R
 - iii) A 246E
 - iv) L 286V
 - v) C 410 Y
 - vi) A 260 V
- 30 vii) A 285 V
 - viii) L 392 V

In accordance with another aspect of the invention, are polyclonal antibodies raised to specific predicted sequences of the ARMP protein. Polypeptides of at least six amino acid residues are provided. The polypeptides of six or greater amino acid residues may define antigenic epitopes of the ARMP. Monoclonal antibodies having suitably specific binding affinity for the antigenic regions of the ARMP are prepared by use of corresponding hybridoma cell lines. In addition, other polyclonal antibodies may be prepared by inoculation of animals with suitable peptides or holoprotein which add suitable specific binding affinities for antigenic regions of the ARMP.

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In accordance with another aspect of the invention, an isolated DNA molecule is provided which codes for E5-1 protein. A plasmid including this nucleic acid was deposited with the ATCC under the terms of the Budapest Treaty on June 28, 1995 and has been assigned ATCC accession number 97214.

In accordance with another aspect of the invention, purified E5-1 protein is provided, having amino acid SEQ ID NO:138.

In accordance with another aspect of the invention a bioassay is provided for determining if a subject has a normal or mutant ARMP, where the bioassay comprises

providing a biological sample from the subject

conducting a biological assay on the sample to detect a normal or mutant gene sequence coding form ARMP, a normal or mutant ARMP amino acid sequence, or a normal or defective protein function.

In accordance with another aspect of the invention, a process is provided for producing ARMP comprising culturing one of the above described transfected host cells under suitable conditions, to produce the ARMP by expressing the DNA sequence. Alternatively, ARMP may be isolated from mammalian cells in which the ARMP normally expressed.

In accordance with another aspect of the invention, a therapeutic composition comprising ARMP and a pharmaceutically acceptable carrier.

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In accordance with another aspect of the invention, a recombinant vector for transforming a mammalian tissue cell to express therapeutically effective amounts of ARMP in the cells is provided. The vector is normally delivered to the cells by a suitable vehicle. Suitable vehicles include vaccinia virus, adenovirus, liposome transport, virus, retrovirus, associated neuraltripic viruses, Herpes simplex virus and other vector systems.

In accordance with another aspect of the invention, a method of treating a patient deficient in normal ARMP the patient comprising administering to therapeutically effective amount of the protein targeted at a variety of patient cells which normally express The extent of administration of normal ARMP being sufficient to override any effect the presence of the mutant ARMP may have on the patient. As an alternative to protein, suitable ligands and therapeutic agents such as small molecules and other drug agents may be suitable for drug therapy designed to replace the protein and 30

defective ARMP, displace mutant ARMP, or to suppress its formation.

another aspect of the with In accordance immuno therapy for treating a patient invention an having Alzheimer's Disease comprises treating the patient with antibodies specific to the ARMP mutant activity or the levels biological reduce To facilitate such amino mutant ARMP in the patient. acid therapy, a vaccine composition may be provided for evoking an immune response in a patient of Alzheimer's disease where the composition comprises a mutant ARMP pharmaceutically acceptable carrier with without a suitable excipient. The antibodies developed specific to the mutant ARMP could be used to target appropriately encapsulated drugs/molecules, specific Therapies utilizing specific sites. cellular/tissue ligands which bind to normal or wild type ARMP of either mutant or wild type and which augments normal function of ARMP in membranes and/or cells or inhibits the deleterious effect of the mutant protein are also made possible.

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In accordance with another aspect of the invention, a transgenic animal model for Alzheimer's Disease which has the mammalian polynucleotide sequence with at least one mutation which when expressed results in mutant ARMP in animal cells and thereby manifests a phenotype. For example, the human Prion gene when overexpressed in rodent peripheral nervous system and muscle cells causes a quite different response in the animal than the human. The animal may be a rodent and is preferably a mouse, but may also be other animals including rat, pig,

Irosophila melanogaster, C. elegans (nematode), all of which are used for transgenic models. Yeast cells can also be used in which the ARMP Sequence is expressed from an artificial vector.

In accordance with another aspect of the invention, a transgenic mouse model for Alzheimer's Disease has the mouse gene encoding ARMP human or murine homologues mutated to manifest the symptoms. The transgenic mouse may exhibit symptoms of cognitive memory or behavioral disturbances. In addition or alternatively, the symptoms may appear as another cellular tissue disorder such as in mouse liver, kidney, spleen or bone marrow or other organs in which the ARMP gene is normally expressed.

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In accordance with another aspect of the invention, the protein can be used as a starting point for rationale drug design to provide ligands, therapeutic drugs or other types of small chemical molecules.

BRIEF DESCRIPTION OF THE DRAWINGS

Various aspects of the invention are described hereinafter with respect to the drawings wherein:

Figure 1a. Genomic physical and transcriptional map of the AD3 region of chromosome 14. Genetic map inter-marker genetic distances averaged for male and female meiosis are indicated in centiMorgans.

Figure 1b. Is the constructed physical contig map of overlapping genomic DNA fragments cloned into YACs spanning a FAD locus on chromosome 14q.

Figure 1c. Regions of interest within the 30 constructed physical contig map.

Figure 1d. Transcriptional map illustrating physical locations of the 19 independent longer cDNA clones.

Figure 2. Automated florescent chromatograms representative the change in nucleic acids which direct (by the codon) the amino acid sequence of the gene.

(a) Met 146 Leu
(b) His 163 Arg
(c) Ala 246 Glu
(d) Leu 286 Val
(e) Cys 410 Tyr

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Figure 3(a). Restriction fragments of M 146 L mutation using BsphI restriction enzyme in AD patients. Absence of a mestriction site indicates a mutant allele.

Figure 3(b). Presence of the His 163 Arg mutation detected by NlaIII restriction digestion.

Absence of a restriction indicates a mutant allele.

Figure 3(c) Presence of the Ala 246 Glu mutation in AD patients using DdeI restriction enzyme. Presence of the mutant allele leads to restriction.

Figure 3(d). Presence of Cys 410 Tyr mutation in AD patients as assayed using allelle specific oligonucleotides.

Figure 3(e). Presence of Leu286Val mutation in AD patients using PvuII restriction enzyme in AD patients.

Figure 4. RNA blot demonstrating the expression of ARMP protein mRNA in different regions of the brain including amygdala, caudate, corpus callosum, hippocampus, hypothalamus, substantia nigra, subthalamic nucleus and thalamus.

Figure 5. RNA blot demonstrating the expression of ARMP protein RNA in a variety of tissues including heart, brain, pracenta, lung, liver, skeletal muscle, kidney and pancreas.

Figure 6a. Hydropathy plot of the putative ARMP protein.

Figure 6b. for the structural model Α Roman of the putative ARMP protein. organization numerals depict the transmembrane domains. Putative glycosylation sites are indicated as asterisks and most 10 of the phosphorylation sites are located on the same membrane face as the two acidic hydrophillic loops. MAP kinase site is present at residue 115 and the PKC site at residue 114. \ FAD mutation sites are indicated by horizontal arrows. 15

investigated \by hybridization of the E5-1 cDNA Northern blots of mRNA from multiple human brain regions (Panel A), and several peripheral tissues (Panel C). brain, the E5-1 thanscript is of lower molecular a lester that the ARMP abundance weight and transcript (Panel B)

hybridized to the same blot using identical conditions.

Figure 8 shows the predicted structure of the E5-1 protein.

DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS

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In order to facilitate review of the various embodiments of the invention and an understanding of various elements and constituents used in making the invention and using same, the following definition of terms used in the invention description is as follows:

Alzheimer Related Membrane Protein gene (ARMP gene) mutated when gene which 14 chromosome Disease and/or Alzheimer's associated with familial other inheritable disease phenotypes (e.g., cerebral schizophrenia, retardation, mental hemorrhage, definition This depression). psychosis, and understood to include the various sequence polymorphisms that exist, wherein nucleotide substitutions in the gene sequence do not affect the essential function of the gene product, as well as functional equivalents of the nucleotide sequences of SEQ ID NO:1, SEQ ID NO:133, SEQ ID NO:3 and SEQ ID NO:135. This term primarily relates to an isolated coding sequence, but can include some or all of the flanking regulatory elements and/or introns. The term ARMP gene includes the gene in other species analogous to the human gene which when mutated is associated with Alzheimer's Disease.

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Alzheimer Related Membrane Protein (ARMP) - the protein encoded by the ARMP gene. The preferred source of protein is the mammalian protein as isolated from Alternatively, functionally animals. humans orequivalent proteins may exist in plants, insects and invertebrates (such as C.elegans). The protein may be produced by recombinant organisms, or chemically is This definition enzymatically synthesized. understood to include functional variants such as the various polymorphic forms of the protein wherein amino acid substitutions or deletions within the amino acid sequence do not affect the essential functioning of the protein, or its structure. It also includes functional fragments of ARMP.

Mutant ARMP gene - The ARMP gene containing one or more mutations which lead to Alzheimer's Disease and/or other inheritable disease phenotypes (e.g., retardation, schizophrenia, mental hemorrhage, This definition psychosis, and depression). understood to include the various mutations that exist, wherein nucleotide substitutions in the gene sequence affect the essential function of the gene product, well as mutations of functional equivalents of the nucleotide sequences of SEQ ID NO:1, SEQ ID NO:133, SEQ ID NO:3 and SEQ ID NO:135 (the corresponding amino acid This term primarily relates to an isolated sequences). coding sequence, but also can include some or all of the flanking regulatory elements and/or introns.

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Mutant ARMP - a mammalian protein that is highly 15 analogous to ARMP in terms of primary structure, but deletions amino acid and/or more wherein one or substitutions result in impairment of its essential function, so that mammals, especially humans, whose ARMP producing cells express mutant ARMP rather than the 20 normal ARMP, demonstrate the symptoms of Alzheimer's Disease and/or other relevant inheritable phenotypes hemorrhage, mental retardation, (e.g. cerebral schizophrenia, psychosis, and depression).

mARMP gene - mouse gene analogous to the human ARMP Functional equivalent as used in describing gene sequences and amino acid sequences means that a recited identical the definitive be to sequence need not sequence of the Sequence ID Nos but need only provide a sequence which functions biologically and/or chemically 30 the equivalent of the definitive sequence.

sequences which correspond to a definitive sequence may also be considered as functionally equivalent sequence.

mARMP - mouse Alzheimer related membrane protein, analogous to the human ARMP, encoded by the mARMP gene. This definition is understood to include the various polymorphic forms of the protein wherein amino acid substitutions or deletions of the sequence does not affect the essential functioning of the protein, or its structure.

Mutant mARMP - a mouse protein which is highly analogous to mARMP in terms of primary structure, acid deletions amino more but wherein one or impairment its substitutions result in and/or essential function, so that mice, whose mARMP producing cells express mutant mARMP rather than the normal mARMP 15 demonstrate the symptoms of Alzheimer's Disease and/or inheritable phenotypes, other other relevant phenotypes and behaviours as manifested in mice.

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ARMP carrier - a mammal in apparent good health whose chromosomes contain a mutant ARMP gene that may be transmitted to the offspring and who will develop Alzheimer's Disease in mid to late adult life.

Missense mutation - A mutation of nucleic acid sequence which alters a codon to that of another amino acid, causing an altered translation product to be made.

Pedigree - In human genetics, a diagram showing the ancestral relationships and transmission of genetic traits over several generations in a family.

E5-1 gene - the chromosome 1 gene which shows homology to the ARMP gene and which when mutated is 30 associated with familial Alzheimer's Disease

other inheritable disease phenotypes. This definition the various sequence understood to include wherein nucleotide exist, that polymorphisms substitutions in the gene sequence do not affect the essential function of the gene product, as well functional equivalents of the nucleotide SEQ ID NO:137. This term also includes the gene in other species analogous to the human gene described herein.

gene. This term includes the protein of SEQ ID NO:138 and also functional variants such as the various polymorphic and splice variant forms of the protein wherein amino acid substitutions or deletions within the amino acid sequence do not affect the essential functioning of the protein. The term also includes functional fragments of the protein.

Mutant E5-1 gene - the E5-1 gene containing one or more mutations which lead to Alzheimer's Disease. This term is understood to include the various mutations that exist, wherein nucleotide substitutions in the gene sequence affect the essential function of the gene product.

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Mutant E5-1 protein - a protein analogous to E5-1 protein but wherein one or more amino acid deletions and/or substitutions result in impairment of its essential function such that mammals, especially humans, whose E5-1 producing cells express mutantE5-1 protein demonstrate the symptoms of Alzheimer's disease.

Linkage analysis-Analysis of co-segregation of a disease trait or disease gene with polymorphic genetic markers of defined chromosomal location.

hARMP gene - Human ARMP gene.

ORF - Open reading frame.

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PCR - Polymerase chain reaction.

contig - continuous cloned regions.

YAC - yeast artificial chromosome.

RT-PCR - reverse transcription polymerase chair 10 reaction.

SSR - Simple sequence repeat polymorphism.

The invention is concerned with present the identification and sequencing of the mammalian ARMP gene in order to gain insight into the cause and etiology of familial Alzheimer's Disease. From this information, 15 screening methods and therapies for the diagnosis and treatment of the disease can be developed. The gene has identified. CDNA isolated and cloned. its been transcripts and gene products identified and sequenced. During such identification of the gene, considerable 20 sequence information has also been developed on intron information in the ARMP gene, flanking untranslated information and information information and signal involving neighbouring genes in the AD3 chromosome 25 region. Direct sequencing of overlapping products spanning the human gene isolated from affected members of large pedigrees linked to chromosome 14 has led to the discovery of missense mutation which cosegregate with the disease.

Although it is generally understood that Alzheimer's Disease is a neurological disorder, most

likely in the brain, expression of ARMP has varieties of human tissue such as heart, found in brain, placenta, lung, liver, skeletal muscle, expressed gene is this and pancreas. Although widely, the clinically apparent phenotype exists is conceivable that biochemical it although in these other tissues. exist may phenotypes as Huntington's diseases such genetic other the clinical Alzheimer's, Disease and APP reflect different biochemistries manifestation may and tissues (which types cell ofdifferent genetics and the protein). Such findings from may not be solely a neurological that ADsuggest disorder but may also be a systemic disorder, hence requiring alternative therapeutic strategies which may be targeted to other tissues or organs or generally in addition or separately from neuronal or brain tissues.

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The ARMP mutations identified have been related to Alzheimer's Disease pathology. With the identification of sequencing of the gene and the gene product, probes and antibodies raised to the gene product can be used in a variety of hybridization and immunological assays to screen for and detect the presence of either a normal or mutated gene or gene product.

patient therapy through removal or blocking of the mutant gene product, as well as supplementation with the normal. gene product by amplification, by genetic and recombinant techniques or by immunotherapy can now be achieved. Correction or modification of the defective gene product by protein treatment immunotherapy (using antibodies to the defective protein) or knock-out of the

mutated gene is now also possible. Familial Alzheimer's Disease could also be controlled by gene therapy in which the gene defect is corrected in situ or by the use of recombinant or other vehicles to deliver a DNA sequence capable of expressing the normal gene product, or a deliberately mutated version of the gene product whose effect counter balances the deleterious consequences of the disease mutation to the affected cells of the patient.

The present invention is also concerned with the identification and sequencing of a second gene, the E5-1 gene on chromosome 1, which is associated with familial Alzheimer's Disease.

Disease mechanism insights and therapies analogous to those described above in relation to the ARMP gene will be available as a result of the identification and isolation of the E5-1 gene.

Isolating the Human ARMP Gene

20 Genetic mapping of the AD3 locus.

After the initial regional mapping of the AD3 gene locus to 14q24.3 near the anonymous microsatellite markers D14S43 and D14S53 (Schellenberg, GD et al., St. George-Hyslop, al., 1992; ₽ et 25 1992; Broeckhoven, C et al., 1992), twenty one pedigrees were used to segregate AD as a putative autosomal dominant trait (St. George-Hyslop P et al., 1992) and to investigate the segregation of 18 additional genetic markers from the 14q24.3 region which had been organized 30 into a high density genetic linkage map (Figure 1b)

1992; Gyapay et al., al., (Weissenbach et previously analyses likelihood Pairwise maximum published confirmed substantial cumulative evidence for linkage between FAD and all of these markers (Table 1). However, much of the genetic data supporting linkage to these markers were derived from six large early onset pedigrees FAD1 (Nee et al., 1983) FAD2 (Frommelt et al., 1991), FAD3 (Goudsmit et al., 1981; Pollen, 1993), FAD4 (Foncin et al., 1985) TOR1.1 (Bergamini, 1991) and 603 (Pericak-Vance et al., 1988) each of which provide at lease one anonymous genetic marker from 14q24.3 (St. George-Hyslop, P. et al., 1992).

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In order to more precisely define the location of the AD3 gene relative to the known locations of the genetic markers from 14q24.3, recombinational landmarks were sought by direct inspection of the raw haplotype data only from genotyped affected members of the six pedigrees showing definitive linkage to chromosome 14. This selective strategy in this particular instance reconstructed from the necessarily discards data genotypes of deceased affected members as well as from elderly asymptomatic members of large pedigrees, and takes no account of the smaller pedigrees of uncertain linkage status. However, this strategy is very sound because it also avoids the acquisition of potentially misleading genotype data acquired either through errors the reconstructed genotypes of deceased affected in members arising from non-paternity or sampling errors or from the inclusion of unlinked pedigrees.

30 Upon inspection of the haplotype data for affected subjects, members of the six large pedigrees whose

genotypes were directly determined revealed obligate recombinants at D14S48 and D14S53, and at D14S258 and at D14S53, which recombinant The single D14S63. the FAD region, depicts a telomeric boundary for the same AD affected of subject occurred in previously been found to be had pedigree who FAD1 recombinant at several other markers located telomeric to D14S53 including D14S48 (St. George-Hyslop, P et al., Conversely, the single recombinant at D14S258, which marks a centromeric boundary of the FAD region, occurred in an affected member of the FAD3 pedigree who several other markers at recombinant also was Both including D14S63. centromeric D14S258 to unequivocal evidence recombinant subjects had Alzheimer's Disease confirmed though standard clinical tests for the illness in other affected members of their families, and the genotypes of both recombinant subjects informative and co-segregating at multiple loci was within the interval centromeric to D14S53 and telomeric to D14S258.

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haplotype analyses enlarged were When the include the reconstructed genotypes of deceased affected members of the six large pedigrees as well as data from the remaining fifteen pedigrees with probabilities for than 0.95, several additional of less linkage recombinants were detected at one or more marker loci within the interval between D14S53 and D14S258. Thus, detected in the was additional recombinant one reconstructed genotype of a deceased affected member of each of three of the larger FAD pedigrees (FAD1, FAD2 other related families), and eight additional and

recombinants were detected in affected members of five However, while some of these smaller FAD pedigrees. recombinants might have correctly placed the AD3 gene within a more defined target region, we were forced to potentially closer regarded these recombinants" as unreliable not only of the reasons they provided earlier, also because but discussed mutually inconsistent locations for the AD3 gene within the D14S53-D14S258 interval.

10 Construction of a Physical Contig Spanning the AD3 Region.

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step toward cloning the AD3 initial As an a contig of overlapping genomic DNA fragments artificial chromosome vectors, into yeast cloned 15 phage artificial chromosome vectors and cosmid vectors was constructed (Figure 1b). FISH mapping the YAC clones from 932c7 derived using cosmids interval most 964f5 suggested that the and at least five carry the gene was AD3 20 large size of Because the in size. megabases region would make minimal co-segregating this cloning strategies intactable, additional positional genetic pointers were sought which focused the search for the AD3 gene to one or more subregions within the 25 interval flanked by D14S53 and D14S258. Haplotype the markers between D14S53 analyses at failed to detect statistically significant evidence for disequilibrium allelic association and/or linkage between the FAD trait and alleles at any of these 30 irrespective of whether the analyses markers, restricted to those pedigrees with early onset forms of FAD, or were generalized to include all pedigrees. result was not unexpected given the diverse ethnic However, when pedigrees of origins of our pedigrees. similar ethnic descent were collated, direct inspection on the disease haplotypes observed the different pedigrees segregating in chromosomes similar ethnic origin revealed two clusters of marker The first of these clusters located loci (Table 2). centromeric to D14S77 (D14S786, D14S277 and D14S268) and spanned the 0.95 Mb physical interval contained in YAC 78842 (depicted as region B in figure 1c). The second telomeric to D14S77 (D14S43, located cluster was 1Mb physical and D14S76) and spanned the -D14S273. included within the overlapping YAC clones interval 964c2, 74163, 797dll and part of 854f5 (depicted as region A in figure 1c). Identical alleles were observed in at least two pedigrees from the same ethnic origin As part of the strategy, it was reasoned (Table 2). that the presence of shared alleles at one of these groups of physically clustered marker loci might reflect physical region co-inheritance of small the founder original on the surrounding the ARMP gene chromosome in each ethnic population. Significantly, each of the shared extended haplotypes were rare in normal caucasian populations and allele sharing was not observed at other groups of markers spanning similar genetic intervals elsewhere on chromosome 14q24.3. Transcription mapping and preliminary analysis of

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candidate genes

To isolate expressed sequences encoded within both critical intervals, a direct selection strategy was

cloned, human immobilized, in involving used the hybridization target to recover genomic DNA as from primary complementary sequences transcribed pools derived from human brain mRNA (Rommens et al., Approximately 900 putative cDNA fragments of size 100 to 600 base pairs were recovered from regions A These fragments were hybridized to and B in figure 1c. Southern blots containing genomic DNAs from each of the overlapping YAC clones and genomic DNAs from humans and This identified a subset of 151 clones other mammals. showed evidence for evolutionary conservation which and/or for a complex structure which suggested that they were derived from spliced mRNA. The clones within this subset were collated on the basis of physical location, cross-hybridization and nucleotide sequence, and were used to screen conventional human brain cDNA At least 19 independent libraries for longer cDNAs. cDNA clones over 1kb in length were isolated and then aligned into a partial transcription map of the AD3 region (Figure 1d). Only three of these transcripts known characterized genes (cFOS, corresponded to latent transferase and dihydrolipoamide succinyl transforming growth factor binding protein 2).

Recovery of Potential Candidate Genes

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Each of the open reading frame portions of the candidate genes were recovered by RT-PCR from mRNA isolated from post-mortem brain tissue of normal control subjects and from either post-mortem brain tissue or cultured fibroblast cell lines of affected members of six pedigrees definitively linked to chromosome 14. The RT-PCR products were then screened for mutations using

endonuclease restriction and chemical cleavage sequence conformational single-strand fingerprinting polymorphism methods (Saleeba and Cotton, 1993; Liu and 1995), and by direct nucleotide sequencing. With one exception, all of the genes examined, although of interest, were not unique to affected subjects, and did not co-segregate with the disease. The single exception was the candidate gene represented by clone S182 which contained a series of nucleotide changes not which altered the subjects, but observed in normal in affected subjects. predicted amino acid sequence differences were Although nucleotide sequence observed in some of the other genes, most were in the 3' untranslated regions and none were unique to Ad-affected subjects.

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The remaining sequences, a subset of which are mapped in Figure 1b together with additional putative transcriptional sequences not identified in Figure 1c, are identified in the sequence listings as 14 through The SEQ ID NOS:14 to 43 represent neighbouring 43. . genes or fragments of neighbouring genes adjacent the gene or possibly additional coding fragments hARMP arising from alternative splicing of the hARMP. NOS:44-126 and SEQ ID NOS:150-160 represent neighboring fragments containing both and exon genomic Such sequences are useful for creating information. primers, for creating diagnostic tests, creating altered sequences and use of adjacent genomic regulatory models. animal better to create sequences

Characterization of the hARMP Gene

Hypridization of the S182 clone to northern blots identified a transcript expressed widely in many areas of brain\and peripheral tissues as a major 3.0 kb transcript and a minor transcript of 7.0 kb (Figures 4 Although the identity of the ~ 5). transcript is unclear, two observations suggest that the ~ 3.0 kb transcript represents an active product of the Hybridization of the S182 clone to northern blots containing mRNA \from a variety of murine tissues, 10 including brain, dentifies only a single transcript identical in size t δ the ~ 3.0 kb human transcript. All of the longer cDNA clones recovered to date (2.6-2.8 kb), which include both 5' and 3' UTRs and which account for the ~ 3.0 kb band $\delta_{\rm P}$ the northern blot, have mapped 15 exclusively to the same \physical region of chromosome From these experiments the ~ 7.0 kb transcript could represent either a rare alternatively spliced or polyadenylated isoform of the ~ 3.0 transcript or could represent another gene with homplogy to S182. 20

The nucleotide sequence of the major transcript was determined from the consensus of eleven independent independent clones and 3 clones from longer cDNA recovered by standard 5' rapid amplification of homology to other significant ends and bears no genes. The cDNA of the sequenced transcript provided in SEQ ID NO:1 and the predicted amino SEQ ID NO:2. The in provided is sequence acid cDNA sequence of another sequenced human clone also provided as SEQ ID NO:133 and its predicted amino acid sequence is provided in SEQ ID NO:134.

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Analysis of the 5' end of multiple cDNA clones and RT-PCR products as well as corresponding genomic clones indicates that the 5' UTR is contained within at least two exons and that transcription either begins from two different start sites and/or that one of the early 5' untranslated exons is alternatively spliced (Table 6). The longest predicted open reading frame contains 467 amino acids with a small alternatively spliced exon of 4 aminc acids at 25 codons from the putative start codon This putative start codon is the first in (Table 3). phase ATG located 63 bp downstream of a TGA stop codon and lacks a classical Kozak consensus sequences around the first two in-phase ATG sequences (Rogaer et al., in other genes lacking classical Like preparation). 'strong' start codons, the putative 5' UTR of the human transcripts are rich in GC.

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Comparison of the nucleic acid and predicted amino acid sequences with available databases using the BLAST amino acid modest paradigms revealed alignment similarity with the C. elegans sperm integral membrane protein SPE-4 (p=1.5e 25, 24-37% identity over three groups of at least fifty residues) and weaker similarity to portions of several other membrane spanning proteins including mammalian chromogranin A and alpha subunit of mammalian voltage dependent calcium channels (Altschul et al., 1990). This clearly established that they are not the same gene. The amino-acid sequence similarities across putative transmembrane domains may occasionally yield alignment that simply arises from the limited number of hydrophobic amino acids, but there is also extended sequence alignment between S182 protein and

hydrophillic Both domains. at several SPE-4 putative S182 protein and SPE-4 are predicted to be of comparable size (467 and 465 residues, respectively) and to contain at least seven transmembrane domains with a predicted final domain preceding the acidic domain acidic domains with a large transmembrane preceding the final predicted transmembrane domain. S182 protein does have a longer predicted hydrophillic region at the N terminus.

hARMP has of the investigation Further 10 form sequence fragments which host οf revealed a intron sequence include and the hARMP gene information, 5' end untranslated sequence information and 3' end untranslated sequence information (Table 6). Such sequence fragments are identified in Sequence ID 15 Nos. 6 to 13.

Mutations in the S182 Transcript

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Direct sequencing of overlapping RT-PCR products kb S182 transcript isolated from spanning the 3.0 affected members of the six large pedigrees linked to chromosome 14 led to the discovery of eight missense mutations in each of the six pedigrees (Table 7, Figure Each of these mutations co-segregated with the 2). [Figures respective pedigrees in the disease 3(a)(b)(c)(d)(e)], and were absent from 142 unrelated neurologically normal subjects drawn from the ethnic origins as the FAD pedigrees (284 unrelated chromosomes).

The location of the gene within the physical interval segregating with AD3 trait, the presence of eight different missense mutations which co-segregate

with the disease train in six pedigrees definitively linked to chromosome 14, and the absence of these independent normal chromosomes 284 in mutations cumulatively confirms that the hARMP gene is the AD3 Further biologic support for this hypothesis arises from the fact that the residues mutated in FAD kindreds are conserved in evolution (Table 3) and occur the protein which are also domains of conserved, and from the fact that the S182 gene product is expressed at high levels in most regions of the brain including the most severely affected with AD.

The DNA sequence for the hARMP gene as cloned has been incorporated into a plasmid Bluescript. This stable vector has been deposited at ATCC under accession number 97124 on April 28, 1995.

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Several mutations in the hARMP gene have been identified which cause a severe type of familial Alzheimer's Disease. One, or a combination of these mutations may be responsible for this form of several Alzheimer's Disease well other as as neurological disorders. The mutations may be any form of nucleotide sequence alteration or substitution. the form of Specific disease causing mutations in nucleotide and/or amino acid substitutions have been located, although we anticipate additional mutations will be found in other families. Each nucleotide substitutions occurred within the putative ORF of the S182 transcript, and would be predicted to acid the following encoded amino at change the positions, numbering from the first putative initiation 30 The mutations are listed in respect of their codon.

nucleotide locations in SEQ ID NO:1 and SEQ ID NO:133 (an additional human clone) and amino acid locations in SEQ ID NO:2 and SEQ ID NO:134 (the additional human clone).

5	i)	685, A→C	Met	146	Leu
	ii)	737, A→G	His	163	Arg
	iii)	986, C→A	Ala	246	Glu
	iv)	1105, C→G	Leu	286	Val
	v)	1478,G→A	Cys	410	Tyr
10	vi)	1027, C→T	Ala	260	Val
	vii)	1102, C→T	Ala	285	Val
	viii) 1422, C→G	Leu	392	Val

The Met146Leu, Ala246Glu and Cys410Tyr mutations have not been detected in the genomic DNA of affected members of the eight remaining small early onset 15 autosomal dominant FAD pedigrees or six additional families in our collection which express late FAD onset. We predict that such mutations would not commonly occur in late onset FAD which has been excluded by genetic linkage studies from the more aggressive form of AD 20 linked to chromosome 14q24.3 (St. George-Hyslop, P. et al., 1992; Schellenberg et al., 1993). The Hisl63Arg mutation has been found in the genomic DNA of affected one additional FAD pedigree for which members of positive but significant statistical evidence 25 linkage to 14 becomes established. Age of onset of affected. consistent with affected members was individuals from families linked to chromosome 14.

Mutations Ala260Val, Ala285Val, and Leu392Val all occur within the acidic hydrophilic loop between putative transmembrane 6 (TM6) and transmembrane (TM7)

(Figure 6). Two of the mutations (A260V; A285V) and the L286V mutation are also located in the alternative spliced domain.

All eight of the mutations can be assayed by a strategies (direct nucleotide sequencing, variety of ligation polymerase oligos, allele specific RT-PCR products using etc.) RFLPs reaction, SSCP, representing the mature mRNA/cDNA sequence or genomic Allele specific oligos were chosen for assaying For the A260V and the A285V mutations, the mutations. genomic DNA carrying the exon was amplified using the same PCR primers and methods as for the L286V mutation. PCR products were then denatured and slot blotted to duplicate nylon membranes using the slot blot protocol described for the C410T mutation.

nucleotide substitutions coof the respective disease in their with the segregated seen (F****gures 3a to 3e), none were in pedigrees asymptomatic family members aged more than two standard deviations beyond the mean age of onset, and non were 20 present on 284 chromosomes from unrelated neurologically normal subjects drawn from comparable ethnic origins.

Identification of an Alternative Splice Form of the ARMP Gene Product

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During sequencing studies of RT-PCR products for the ARMP gene recovered from a variety of tissues, it was discovered that some peripheral tissues (principally white blood cells) demonstrated two alternative splice forms of the ARMP gene. One form is identical to the (putatively 467 amino acid) isoform constitutatively

expressed in all brain regions. The alternative splice

form results from the exclusion of the segment of the cDNA between base pairs 1018 and 1116 inclusive, and results in a truncated isoform of the ARMP protein the hydrophilic οf hydrophobic part wherein the acidically-charged loop immediately C-terminal to TM6 is This alternatively spliced isoform therefore is characterized by preservation of the sequence Nterminal to and including the tyrosine at position 256, alanine, changing of the aspartate at 257 to splicing on to the C-terminal part of the protein from Such splicing differences and including tyrosine 291. are often associated with important functional domains of the proteins. This argues that this hydrophilic loop (and consequently the N-terminal hydrophilic loop with amino acid charge) is/are active functional similar and thus sites for the ARMP product domains of therapeutic targeting.

ARMP Protein

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With respect to DNA SEQ ID NO.1 and DNA SEQ NO:133, analysis of the sequence of overlapping cDNA 20 clones predicted an ORF protein of 467 amino acids when read from the first in phase ATG start codon and a approximately 52.6kDa as molecular mass of described, due to either polymorphisms in the protein or alternate splicing of the transcript, the molecular 25 the protein can vary due to possible weight of substitutions or deletions of amino acids.

The analysis of predicted amino acid sequence using the Hopp and Woods algorithm suggested that the protein product is a multispanning integral membrane protein such as a receptor, a channel protein, or a structural

membrane protein. The absence of recognizable signal peptide and the paucity of glycoslyation sites are noteworthy, and the hydropathy profile suggests that the protein is less likely to be a soluble protein with a highly compact three-dimensional structure.

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The protein may be a cellular protein with a highly compact three dimensional structure in which respect is to APOE which is also related may be similar light of this putative In Alzheimer's Disease. functional role, it is proposed that this protein be Alzheimer Related Membrane Protein the labeled as The protein also contains a number of potential (ARMP). phosphorylation sites, one of which is the consensus site for MAPkinase which is also involved in hyperphosphorlyation of tau during the normal conversion to neurofibrillary tangles. This tau normal consensus sequence may provide a common putative pathway linking this protein and other known biochemical aspects of Alzheimer's Disease and would represent a likely therapeutic target. Review of the protein structure reveals two sequence YTPF (residues 115-119) SEQ ID NO:161 and STPE (residues 353-356) SEQ ID NO:162 which represent the 5/T-P motif which is the MAP kinase consensus sequence. Several other phosphorylation sites exist with consensus sequences for Protein Kinase C Because protein kinase C activity activity. associated with differences in the metabolism of APP which are relevant to Alzheimer's Disease, these sites the ARMP protein and homologues are sites for therapeutic targeting.

a highly characterized by N-terminal is The hydrophilic acid charged domain with several potential phosphorylation domains, followed sequentially by hydrophobic membrane spanning domain of 19 residues; a additional five then hydrophilic loop, charged hydrophobic membrane spanning domains interspersed with short (5-20 residue) hydrophilic domains; an additional larger acidic hydrophilic charged loop, and then at least one and possibly two other hydrophobic potentially membrane spanning domains culminating in a polar domain at the C-terminus (Table 4 and Figure 6B). The presence of seven membrane spanning domains is characteristic of several classes of G-coupled receptor proteins but is also observed with other proteins including channel proteins.

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Comparison of the nucleic acid and predicted amino acid sequences with available databases using the BLAST alignment paradigms revealed amino acid similarity with the C. elegans sperm integral membrane protein spe-4 and a similarity to several other membrane spanning proteins including mammalian chromogranin A and the α -subunit of mammalian voltage dependent calcium channels.

The similarity between the putative products of the spe-4 and ARMP genes implies that they may have similar activities. The SPE-4 protein of C. elegans appears to be involved in the formation and stabilization of the fibrous body-membrane organelle (FMBO) complex during spermatogenesis. The FBMO is a specialized Golgiderived organelle, consisting of a membrane bound vesicle attached to and partly surrounding a complex of parallel protein fibers and may be involved in the

transport and storage of soluble and membrane-bound Mutations in spe-4 disrupt the polypeptides. complexes and arrest spermatogenesis. Therefore the physiologic function of spe-4 may be either to stabilize integral membrane budding interactions between fusion events, or to stabilize interactions between the proteins during the fibrillary membrane and complex during transport of the FBMO intracellular functions could be Comparable spermatogenesis. The ARMP could be involved envisaged for the ARMP. either in the docking of other membrane-bound proteins such as etaAPP, or the axonal transport and fusion budding of membrane-bound vesicles during protein transport such as in the golgi apparatus or endosome-lysosome system. If correct, then mutations might be expected to result in aberrant transport and processing of etaAPP and/or abnormal interactions with cytoskeletal proteins such as the microtubule-associated protein Tau. Abnormalities extracellular intracellular in the and in the disposition of both etaAPP and Tau are in fact an integral the neuropathologic features of Alzheimer's Although the location of the ARMP mutations in Disease. highly conserved residues within conserved domains of the putative proteins suggests that they are pathogenic, at least three of these mutations are conservative which is commensurate with the onset of disease in adult life. Because none of the mutations observed so far are deletions or nonsense mutations that would be expected to cause a loss of function, we cannot predict whether these mutations will have a dominant gain-of-function effect and promote aberrant processing of etaAPP or a

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dominant loss-of-function effect causing arrest of normal etaAPP processing.

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An alternative possibility is that the ARMP gene product may represent a receptor or channel protein. Mutations of such proteins have been causally related to several other dominant neurologic disorders in both vertebrate (e.g., Malignant hyperthermia, hyperkalemic in invertebrate and humans) periodic paralysis in organisms (deg-1(d) mutants in C. elegans). the pathology of these other disorders does not resemble is evidence that of Alzheimer's Disease there functional abnormalities in ion channels in Alzheimer's For example, anomalies have been reported in Disease. tetra-ethylammonium-sensitive 113pS Perturbations in channel and in calcium homeostasis. especially calcium fluxes might be transmembrane relevant in view of the weak homology between S182 and the $\alpha\text{-ID}$ subunit of voltage-dependent calcium channels and the observations that increases in intracellular calcium in cultured cells can replicate some of the biochemical features of Alzheimer's Disease such as alteration in the phosphorylation of Tau-microtubuleassociated protein and increased production of ABpeptides.

protein mentioned purified normal ARMP is characterized by a molecular weight of 52.6kDa. The substantially free of protein, ARMP normal proteins, is encoded by the aforementioned SEQ ID NO:.1 and SEQ ID NO:133. As will be later discussed, the ARMP protein and fragments thereof may be made by a variety is ARMP protein methods. Purified mutant of

characterized by FAD-associated phenotype apoptic death, granulovascular degeneration, neurofibrillary degeneration, abnormalities or changes in the metabolism of APP, and Ca^{2+} , K^{+} and glucose, and metabolism energy and function mitochondrial which have been neurotransmitter metabolism, all of found to be abnormal in human brain, and/or peripheral tissue cells in subjects with Alzheimer's Disease) in a The mutant ARMP, free of other variety of cells. proteins, is encoded by the mutant DNA sequence.

Description of the E5-1 gene, a Homologue of the ARMP Gene

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A gene, E5-1, with substantial nucleotide and amino acid homology to the ARMP gene was identified by using the nucleotide sequence of the cDNA for ARMP to search data bases using the BLASTN paradigm of Atschul et al., 1990. Three expressed sequence tagged sites (ESTs) identified by accession numbers T03796, R14600, and R05907 were located which had substantial homology (p < $1.0 \, {\rm e}^{-100}$, greater than 97% identity over at least 100 contiguous base pairs).

Oligonucleotide primers were produced from these sequences and used to generate PCR products by reverse transcriptase PCR (RT-PCR). These short RT-PCR products were partially sequenced to confirm their identity with the sequences within the data base and were then used as full-length CDNA screen to probes hybridization Several different cDNA's ranging in size libraries. from 1Kb to 2.3Kb were recovered from a cancer cell cDNA library (CaCo-2) and from a human brain cDNA library (E5-1, G1-1, cc54, cc32).

The nucleotide sequence of these clones confirmed that all were derivatives of the same transcript (designated E5-1). A plasmid including this nucleic acid was deposited with the ATCC under the terms of the Budapest Treaty on June 28, 1995 and has been assigned ATCC accession number 97214.

The gene encoding the E5-1 transcript mapped to human chromosome 1 using hybrid mapping panels and to two clusters of CEPH Mega YAC clones which have been placed upon a physical contig map (YAC clones 750g7, 921d12 mapped by FISH to 1q41; and YAC clone 787g12 which also contains an EST of the leukemia associated phosphoproteins (LAP18) gene which has been mapped to 1p36.1-p35) (data not shown).

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Hybridization of the E5-1 cDNA clones to Northern Blots detected an ~2.3 kilobase mRNA band in many tissues including regions of the brain, as well as a ~2,6K.b mRNA band in muscle, cardiac muscle and pancreas (Figure 7).

In skeletal muscle, cardiac muscle and pancreas, the E5-1 gene is expressed at relatively higher levels than in brain and as two different transcripts of ~2.3Kb and ~2.6Kb. Both of the E5-1 transcripts have sizes clearly distinguishable from that of the 2.7Kb ARMP transcript, and did not cross-hybridize with ARMP probes at high stringency. The cDNA sequence of the E5-1 gene is identified as SEQ ID NO.:137.

The longest ORF within the E5-1 cDNA consensus nucleotide sequence predicts a polypeptide containing 448 amino acids (numbering from the first in-phase ATG

codon which was surrounded by a <u>GCC</u>-agg-GCt-<u>ATG</u>-c Kozak consensus sequence) (SEQ ID NO.:138).

A comparison of the amino acid sequences of hARMP and E5-1 homologue protein are shown in Table 8. Identical residues are indicated by vertical lines. The locations of mutations in the E5-1 gene are indicated by downward pointing arrows. The locations of the mutations in the hARMP gene are indicated by upward pointing arrows. Putative TM domains are in open ended boxes. The alternatively spliced exons are denoted by superscripted (E5-1) or subscripted (hARMP) "*".

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BLASTP alignment analyses also detected significant (P=3.5e-26;SPE-4 C. elegans of homology with identity=20-63% over five least 22 domains of at residues), and weak homologies to brain sodium channels (alpha III subunit) and to the alpha subunit of voltage dependent calcium channels from a variety of species (P=0.02; identities 20-28% over two or more domains each 35 residues) (Atschul, 1990). of at least alignments are similar to those described above for the ARMP gene. However, the most striking homology to the E5-1 protein was found with the amino acid sequence predicted for ARMP. ARMP and E-51 proteins share 63% acid sequence identity, and amino overall domains display virtually complete identity (Table 8). Furthermore, all eight residues mutated in ARMP subjects with AD3 are conserved in the E5-1 protein would be expected, hydrophobicity As 8). (Table analyses suggest that both proteins also share a similar structural organization.

The similarity was greatest in several domains of the protein corresponding to the intervals between transmembrane domain 1 (TM1) and TM6, and from TM7 to the C-terminus of the ARMP gene. The main difference from ARMP is a difference in the size and amino acid sequence of the acidically-charged hydrophilic loop in the position equivalent to the hydrophilic loop between transmembrane domains TM6 and TM7 in the ARMP protein and in the sequence of the N-terminal hydrophilic domains.

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Thus, both proteins are predicted to possess seven hydrophobic putative transmembrane domains, and both proteins bear large acidic hydrophilic domains at the Nterminus and between TM6 and TM7 (figs. 6 and 8). Α analysis of RT-PCR similarity arose from further products from brain and muscle RNA, which revealed that nucleotides E5-1 transcript 1153-1250 of the These nucleotides encode amino alternatively spliced. acids 263-296, which are located within the TM6-TM7 loop domain of the putative E5-1 protein, and which share 94% the alternatively spliced sequence identity with residues 257-290 in ARMP.

The most noticeable differences between the two predicted amino acid sequences occur in the amino acid of in the central portion the sequence hydrophilic loop (residues 304-374 of ARMP; 301-355 of E5-1), and in the N-terminal hydrophilic domain (Table analogy, this domain is also less highly By between the murine and human ARMP conserved (identity=47/60 residues), and shows no similarity with the equivalent region of SPE-4.

A splice variant of the E5-1 cDNA sequence identified as SEQ ID NO:137 has also been found in all tissues examined. This splice variant lacks the tripet GAA at nucleotide positions 1338-1340.

A further variant has been found in one normal individual whose E5-1 cDNA had C replacing T at nucleotide position 626, which does not change the amino acid sequence.

Mutations of the E5-1 Gene 10 Associated with Alzheimer's Disease

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The strong similarity between ARMP and the E5-1 gene product raised the possibility that the E5-1 gene might be the site of disease-causing mutations in some of a small number of early onset AD pedigrees in which genetic linkage studies have excluded chromosomes 14, 19 and 21. RT-PCR was used to isolate cDNAs corresponding to the E5-1 transcript from lymphoblasts, fibroblasts or post-mortem brain tissue of affected members of eight pedigrees with early onset familial AD (FAD) in which mutations in the β APP and ARMP gene had previously been excluded by direct sequencing studies.

Examination of these RT-PCR products detected a heterozygous A→G substitution at nucleotide 1080 in all four affected members of an extended pedigree of Italian origin (Flo10) with early onset, pathologically confirmed FAD (onset=50-70 yrs.). This mutation would be predicted to cause a Met→Val missense mutation at codon 239 (Table 8).

A second mutation (A→T at nucleotide 787) causing a

Asn→Ile substitution at codon 141 was found in affected

members of a group of related pedigrees of Volga German

ancestry (represented by cell lines AG09369, AG09907, AG09952, and AG09905, Coriell Institute, Camden, NJ). Significantly, one subject (AG09907) was homozygous for this mutation, an observation compatible with the inbred nature of these pedigrees. Significantly, this subject did not have a significantly different clinical picture from those subjects heterozygous for the Arg14Ile mutation. Neither of the E5-1 gene mutations were found in 284 normal Caucasian controls nor were they present in affected members of pedigrees with the AD3 type of AD.

Both of these mutations would be predicted to cause substitutions of residues which are highly conserved within the ARMP/E5-1 gene family.

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The finding of a gene whose product is predicted to substantial amino acid and structural similarities with the ARMP gene product suggest that these proteins may be functionally related either as independent proteins with overlapping functions but perhaps with slightly different specific activities, as multimeric physically associated of а subunits independent proteins performing polypeptide or as consecutive functions in the same pathway.

The observation of two different missense mutations
in conserved domains of the E5-1 protein in subjects
with a familial form of AD argues that these mutations
are, like those in the ARMP gene, causal to AD. This
conclusion is significant because, while the disease
phenotypes associated with mutations in the ARMP gene
(onset 30-50 yrs., duration 10 years) are subtly
different from that associated with mutations in the E5-

1 gene (onset 40-70 years; duration up to 20 years), the general similarities clearly argue that the biochemical pathway subsumed by members of this gene family is central to the genesis of at least early onset AD. The subtle differences in disease phenotype may reflect a lower level of expression of the E5-1 transcript in the CNS, or may reflect a different role for the E5-1 gene product.

By analogy to the effects of ARMP mutations, E5-1 when mutated may cause aberrant processing of APP 10 $A\beta$ peptide, into Protein) Precursor (Amyloid microtubule associated hyperphosphorylation of Tau intracellular calcium abnormalities of and protein with these anomalous Interference homeostasis. interactions provides a potential therapy for AD. 15

Functional Domains of the ARMP Protein are Defined by Splicing Sites and Similarities within Other Members of a Gene Family

The ARMP protein is a member of a novel class of 20 transmembrane proteins which share substantial amino The homology is sufficient that certain acid homology. nucleotides probes and antibodies raised against one can identify other members of this gene family. The major between members of this family 25 difference the amino acid and nucleotide sequence homologous to the hydrophillic acid loop domain between putative and transmembrane 7 domains of transmembrane 6 is gene product. This region and gene tissues, spliced non-neural alternatively in some 30 pathogenic site of several also the is in the ARMP gene. disease-causing mutations

this hydrophilic loop, variable splicing of a high-density of pathogenic mutations presence of the fact that the within this loop, and loop differs between members acid sequences of the family suggest that this loop is the gene functional domain of the protein and important physiologic and specificity to the some pathogenic interactions which the ARMP gene product the N-terminal hydrophilic undergoes because shares the same acidic charge and same orientation with respect to the membrane, it is very likely that these two domains share functionality either in a coordinated independent fashion (e.g., different (together) or ligands or functional properties). As а result everything said about the hydrophilic loop shall apply also to the N-terminal hydrophilic domain.

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Knowledge of the specificity of the loop can be used to identify ligands and functional properties of the ARMP gene product (e.g. sites of interactions with APP, cytosolic proteins such as kinases, Tau, and MAP, Soluble recombinant fusion proteins can be the nucleotide sequence coding for made or within the loop or parts of the loop can be (yeast-2-hybrid, suitable vectors expressed in baculovirus, and phage -display systems for instance), and used to identify other proteins which interact with ARMP in the pathogenesis of Alzheimer's Disease and other neurological and psychiatric diseases. Therapies designated to modulate these interactions and be can thus to modulate Alzheimer's Disease and the other conditions associated with inherited acquired or

abnormalities of the ARMP gene or its gene products. The potential efficacy of these therapies can be tested of by analyzing the affinity function and interactions after exposure to the therapeutic agent by standard pharmacokinetic measurements of affinity (Kd and Vmax etc.) using synthetic peptides or recombinant proteins corresponding to functional domains of the ARMP An alternate method for gene (or its homologues). involving assaying the effect of any interactions as the hydrophilic loop is functional domains such to monitor changes in the intracellular trafficking and post-translational modification of the ARMP gene by immunohistochemistry, Western in-situ hybridization, labeling studies blotting and metabolic pulse-chase absence of the the presence of and in the in monitor therapeutic agents. A third way is to of "downstream" events including (i) effects the intracellular metabolism, changes in the trafficking and targeting of APP and its products; (ii) changes in second messenger event e.g., cAMP, intracellular Ca** protein kinase activities, etc.

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Isolation and Purification of the ARMP Protein

The ARMP protein may be isolated and purified by methods selected on the basis of properties revealed by its sequence. Since the protein possesses properties of a membrane-spanning protein, a membrane fraction of cells in which the protein is highly expressed (e.g., system cells or cells from other nervous tissues) would be isolated and the proteins removed by extraction and the proteins solubilized using detergent.

protein achieved using Purification can be purification procedures such as chromatography methods ion-exchange and immunoaffinity), by (gel-filtration, high-performance liquid chromatography (RP-HPLC, ionsize-exclusion HPLC, high-performance exchange HPLC, interaction hydrophobic chromatofocusing and precipitation by chromatography) or Polyacrylamide (immunoprecipitation). electrophoresis can also be used to isolate the ARMP protein based on its molecular weight, charge properties and hydrophobicity.

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Similar procedures to those just mentioned could be used to purify the protein from cells transfected with vectors containing the ARMP gene (e.g., baculovirus system, yeast expression systems, eukaryotic expression systems).

purified protein can be used in further biochemical analyses to establish secondary and tertiary structure which may aid in the design of pharmaceuticals to interact with the protein, alter protein charge configuration or charge interaction with other proteins, lipid or saccharide moieties, alter its function in membranes as a transporter channel or receptor and/or in cells as an enzyme or structural protein and treat the disease.

The protein can also be purified by creating protein legating the ARMP CDNA by fusion sequence to a vector which contains sequence for (e.g., GST-glutathionine succinyl peptide another The fusion protein is expressed transferase). recovered from prokaryotic (e.g., bacterial or

eukaryotic cells. The fusion baculovirus) or be purified affinity by protein can then vector fusion based upon the chromatography further protein can then be The ARMP sequence. fusion protein by enzymatic the from purified cleavage of the fusion protein.

Isolating Mouse ARMP Gene

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physiological the characterize order to significance of the normal and mutant hARMP gene and in a transgenic mouse model it was gene products necessary to recover a mouse homologue of the hARMP We recovered a murine homologue for the hARMP gene by screening a mouse cDNA library with a labelled human DNA probe and in this manner recovered a 2 kb partial transcript (representing the 3' end of the gene) and several RT-PCR products representing the 5' end. Sequencing of the consensus cDNA transcript of the revealed substantial amino homologue murine The sequence cDNA is identified in SEQ ID identity. NO:3 and the predicted amino acid sequence is provided in SEQ ID NO:4. Further sequencing of the mouse cDNA transcript has provided the sequence of the complete coding sequence identified as SEQ ID NO:135 and the predicted amino acid sequence from this sequence provided in SEQ ID NO:136. More importantly, all of the amino acids that were mutated in the FAD pedigrees were conserved between the murine homologue and the normal human variant (Table 3). This conservation of the ARMP that 3, indicates shown in Table as is gene orthologous gene exists in the mouse (mARMP), and it is now possible to clone mouse genomic libraries using human ARMP probes. This will also make it possible to identify and characterize the ARMP gene in other species. This also provides evidence of animals with various disease states or disorders currently known or yet to be elucidated.

Transgenic Mouse Model

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The creation of a mouse model for Alzheimer's Disease is important to the understanding of the disease and for the testing of possible therapies. Currently no unambiguous viable animal model for Alzheimer's Disease exists.

There are several ways in which to create an animal model for Alzheimer's Disease. Generation of a specific mutation in the mouse gene such as the identified hARMP Secondly, we could gene mutations is one strategy. insert a wild type human gene and/or humanize the murine gene by homologous recombination. Thirdly, it is also possible to insert a mutant (single or multiple) human gene as genomic or minigene cDNA constructs using wild elements. mutant or artificial promoter type or Fourthly, knock-out of the endogenous murine genes may artificially insertion of accomplished by the be modified fragments of the endogenous gene by homologous The modifications include insertion of recombination. mutant stop codons, the deletion of DNA sequences, or the inclusion of recombination elements (lcx p sites) recognized by enzymes such as Cre recombinase.

To inactivate the mARMP gene chemical or x-ray mutagenesis of mouse gametes, followed by fertilization, can be applied. Heterozygous offspring can then be identified by Southern blotting to demonstrate loss of

one allele by dosage, or failure to inherit one parental allele using RFLP markers.

To create a transgenic mouse a mutant version of hRMP or mARMP can be inserted into a mouse germ line using standard techniques of oocyte microinjection or microinjection into stem transfection or Alternatively, if it is desired to inactivate or replace endogenous mARMP gene, homologous recombination using embryonic stem cells may be applied.

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For oocyte injection, one or more copies of the 10 mutant or wild type ARMP gene can be inserted into the pronucleus of a just-fertilized mouse oocyte. This oocyte is then reimplanted into a pseudo-pregnant foster The liveborn mice can then be screened for mother. integrants using analysis of tail DNA for the presence 15 of human ARMP gene sequences. The transgene can be either a complete genomic sequence injected as a YAC, BAC, PAC or other chromosome DNA fragment, a cDNA with either the natural promoter or a heterologous promoter, or a minigene containing all of the coding region and 20 other elements found to be necessary for optimum expression.

Retroviral infection of early embryos can also mutant or wild type hARMP. done to insert the be type hARMP is this method, the mutant or wild inserted into a retroviral vector which is used directly infect mouse embryos during the early stages of development to generate a chimera, some of which will Similar experiments can lead to germline transmission. be conducted in the cause of mutant proteins, using 30 mutant murine or other animal ARMP gene sequences.

Homologous recombination using stem cells allows for screening of gene transfer cells to identify the rare homologous recombination events. Once identified, these can be used to generate chimeras by injection of mouse blastocysts, and a proportion of the resulting transmission from germline mice will show recombinant line. This methodology is especially useful inactivation of the mARMP gene is desired. example, inactivation of the mARMP gene can be done by designing a DNA fragment which contains sequences from a mARMP exon flanking a selectable marker. Homologous recombination leads to the insertion of the marker sequences in the middle of an exon, inactivating the mARMP gene. DNA analysis of individual clones can then be used to recognize the homologous recombination events.

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It is also possible to create mutations in the mouse germline by injecting oligonucleotides containing the mutation of interest and screening the resulting cells by PCR.

This embodiment of the invention has the most significant commercial value as a mouse model for Alzheimer's Disease. Because of the high percentage of sequence conservation between human and mouse it is contemplated that an orthologous gene will exist also in many other species. It is thus contemplated that it will be possible to generate other animal models using similar technology.

Screening and Diagnosis for Alzheimer's Disease

30 General Diagnostic Uses of the ARMP Gene and Gene
Product

The ARMP gene and gene products will be useful for diagnosis of Alzheimer's Disease, presenile and senile dementias, psychiatric diseases such as schizophrenia, and neurologic diseases such depression, etc., hemorrhage - all of which are cerebral and stroke in lesser extent greater or seen to bearing mutations in the ARMP subjects symptomatic Diagnosis of inherited cases gene or in the APP gene. of these diseases can be accomplished by analysis of the cDNA sequence (including genomic and nucleotide sequences included in this patent). Diagnosis can also monitoring alterations in the achieved by by the reaction with electrophoretic mobility and specific antibodies to mutant or wild-type ARMP gene products, and by functional assays demonstrating altered function of the ARMP gene product. In addition, the ARMP gene and ARMP gene products can be used to search for inherited anomalies in the gene and/or its products (as well as those of the homologous gene) and can also be used for diagnosis in the same way as they can be used for diagnosis of non-genetic cases.

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Diagnosis of non-inherited cases can be made by observation of alterations in the ARMP transcription, translation, and post-translational modification and processing as well as alterations in the intracellular and extracellular trafficking of ARMP gene products in the brain and peripheral cells. Such changes will include alterations in the amount of ARMP messenger RNA and/or protein, alteration in phosphorylation state, abnormal intracellular location/distribution, abnormal extracellular distribution, etc. Such assays will

Northern Blots (with ARMP-specific and ARMP include: non-specific nucleotide probes which also cross-react with other members of the gene family), and Western blots and enzyme-linked immunosorbent assays (with antibodies raised specifically to: ARMP; various functional domains of ARMP; to other members of family; and to various homologous gene translational modification states including glycosylated phosphorylated isoforms). These assays can be performed on peripheral tissues (e.g., blood cells, plasma, cultured or other fibroblast tissues, etc.) as well as on biopsies of CNS tissues obtained antimortem or postmortem, and upon cerebrospinal fluid. assays might also include in-situ hybridization and immunohistochemistry (to localized messenger RNA and protein to specific subcellular compartments and/or associated with within neuropathological structures these diseases such as neurofibrillary tangles and amyloid plaques).

Screening for Alzheimer's Disease 20

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Screening for Alzheimer's Disease as linked chromosome 14 may now be readily carried out because of the knowledge of the mutations in the gene.

People with a high risk for Alzheimer's Disease family pedigree) or, individuals (present in previously known to be at risk, or people in general may be screened routinely using probes to detect the present a mutant ARMP gene by a variety of techniques. Genomic DNA used for the diagnosis may be obtained from body cells, such as those present in the blood, tissue 30 biopsy, surgical specimen, or autopsy material. The DNA

may be isolated and used directly for detection of a specific sequence or may be PCR amplified prior to RNA or cDNA may also be used. To detect a analysis. sequence hybridization using specific DNA direct DNA sequencing, restriction oligonucleotides, enzyme digest, RNase protection, chemical cleavage, and ligase-mediated detection are all methods which can be Oligonucleotides specific to mutant sequences can be chemically synthesized and labelled radioactively with isotopes, or non-radioactively using biotin tags, and hybridized to individual DNA samples immobilized on dot-blot solid-supports by other membranes or transfer from gels after electrophoresis. The presence or absence of these mutant sequences are then visualized using methods such as autoradiography, fluorometry, or colorimetric reaction. Examples of suitable PCR primers which are useful for example in amplifying portions of sequence containing the aforementioned subject mutations are set out in Table 5. This table also sets out the change in enzyme site to provide a useful diagnostic tool as defined herein.

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Direct DNA sequencing reveals sequence differences between normal and mutant ARMP DNA. Cloned DNA segments may be used as probes to detect specific DNA segments. PCR can be used to enhance the sensitivity of this method. PCR is an enzymatic amplification directed by sequence-specific primers, and involves repeated cycles of heat denaturation of the DNA, annealing of the complementary primers and extension of the annealed primer with a DNA polymerase. This results in an exponential increase of the target DNA.

Other nucleotide sequence amplification techniques may be used, such as ligation-mediated PCR, anchored PCR and enzymatic amplification as would be understood by those skilled in the art.

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Sequence alterations may also generate fortuitous restriction enzyme recognition sites which are revealed by the use of appropriate enzyme digestion followed by DNA fragments carrying the site gel-blot hybridization. (normal or mutant) are detected by their increase or reduction in size, or by the increase or decrease of corresponding restriction fragment numbers. Genomic DNA samples may also be amplified by PCR prior to treatment the the appropriate restriction and enzyme with fragments of different sizes are visualized under UV light in the presence of ethidium bromide after gel electrophoresis.

Genetic testing based on DNA sequence differences by detection of alteration achieved be may electrophoretic mobility of DNA fragments in gels. and insertions can be deletions sequence Small high resolution gel electrophoresis. visualized by Small deletions may also be detected as changes in the in DNA heteroduplexes οf pattern migration denaturing gel electrophoresis. Alternatively, a single base substitution mutation may be detected based differential PCR product length in PCR. The PCR could be normal and mutant gene the products of differentially detected in acrylamide gels.

Nuclease protection assays (S1 or ligase-mediated)
30 also reveal sequence changes at specific location.

Alternatively, to confirm or detect a polymorphism restriction mapping changes ligated PCR, ASO, REF-SSCP chemical cleavage, endonuclease cleavage at mismatch sites and SSCP may be used. Both REF-SSCP and SSCP are mobility shift assays which are based upon the change in conformation due to mutations.

DNA fragments may also be visualized by methods in which the individual DNA samples are not immobilized on The probe and target sequences may be in solution or the probe sequence may be immobilized. radioactive decay, spectrophotometry, Autoradiography, used to be also may fluorometry mutations Finally, individual genotypes. specific be detected by direct nucleotide sequencing. can

According to an embodiment of the invention, the 15 portion of the cDNA or genomic DNA segment that is informative for a mutation, can be amplified using PCR. For example, the DNA segment immediately surrounding the C410Y mutation acquired from peripheral blood samples screened using the individual can be from an 20 oligonucleotide primers 885 (tggagactggaacacaac) SEQ ID NO:127 and 893 (gtgtggccagggtagagaact) SEQ ID NO:128. This region would then be amplified by PCR, the products to electrophoresis, and transferred by separated Labelled oligonucleotide probes are then 25 membrane. hybridized to the DNA fragments and autoradiography performed.

ARMP Expression

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As an embodiment of the present invention, ARMP 30 protein may be expressed using eukaryotic and prokaryotic expression systems. Eukaryotic expression

systems can be used for many studies of the ARMP gene and gene product including determination of proper expression and post-translational modifications for full biological activity, identifying regulatory elements located in the 5' region of the ARMP gene and their role in tissue regulation of protein expression, production of large amounts of the normal and mutant protein for isolation and purification, to use cells expressing the ARMP protein as a functional assay system for antibodies generated against the protein or to test effectiveness of pharmacological agents, or as a component of a signal transduction system, to study the function of the normal complete protein, specific portions of the protein, or of naturally occurring and artificially produced mutant proteins.

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Eukaryotic and prokaryotic expression systems were generated using two different of ARMP classes the first sequence inserts. In CDNA nucleotide the constructs, full-length termed class. the into is inserted sequence cDNA 20 ARMP expression plasmid in the correct orientation, and 5' UTR and both the natural includes entire open the as sequences as well nucleotide frames bear a open reading The frame. the wild which allows either cassette 25 sequence in included reading frame be to open system or alternatively, a single expression combination of double mutations can be inserted into the This was accomplished by removing a open reading frame. restriction fragment from the wild type open reading 30 frame using the enzymes NarI and PflmI and replacing it

generated by fragment similar with а transcriptase PCR which bears the nucleotide sequence encoding either the Met146Leu mutation or the Hys163Arg A second restriction fragment was removed mutation. from the wild type normal nucleotide sequence for the open reading frame by cleavage with the enzymes PflmI and NcoI and replaced with restriction fragments bearing Ala246Glu the nucleotide sequence encoding mutation, or the Ala260Val mutation or the Ala285Val mutation or the Leu286Val mutation, or the Leu392Val Finally, a third mutation, or the Cys410Tyr mutation. variant bearing combinations of either the Met146Leu or remaining in tandem with the mutations His163Arq mutations, was made by linking the Narl-PflmI fragment bearing these mutations and the PflmI-NcoI fragments bearing the remaining mutations.

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above.

A second variant of cDNA inserts bearing wild type or mutant cDNA sequences was constructed by removing from the full-length cDNA the 5' UTR and part of the 3' UTR sequences. The 5' UTR sequence was replaced with a synthetic oligonucleotide containing a KpnI restriction site and a Kozak initiation site (oligonucleotide 969: ggtaccgccaccatgacagaggtacctgcac, SEQ ID NO:139). UTR was replaced with an oligonucleotide corresponding to position 2566 of the cDNA and bears an artificial (oligonucleotide 970: site ECORI NO:140). Mutant SEQ ID gaattcactggctgtagaaaaagac, variants of this construct were then made by inserting the same mutant sequences described above at the NarI-PflmI fragment, and at the PsImI-NcoI sites described

eukaryotic expressions, these various type and mutant sequences bearing wild constructs described above were cloned into the expression vector pZeoSV (invitrogen). For prokaryotic expression, using the glutathione constructs have been made The inserts vector pGEX-kg. fusion transferase have been attached GST fusion to the which nucleotide same nucleotide sequence are the (generated the with sequence described above ID NO:139 and 970, 969, SEQ oligonucleotide primers SEQ ID NO:140) bearing either the normal open reading frame nucleotide sequence, or bearing a combination of single and double mutations as described above. This construct allows expression of the full-length protein in mutant and wild type variants in prokaryotic cell which allows GST fusion protein as a systems purification of the full length protein followed by removal of the GST fusion product by thrombin digestion. The second prokaryotic cDNA construct was generated to create a fusion protein with the same vector, and allows the production of the amino acid sequence corresponding to the hydrophilic acidic loop domain between TM6 and TM7 of the full-length protein, as either a wild type sequence (thus a wild type amino nucleotide sequence for fusion proteins) or as a mutant sequence bearing either the Ala285Val mutation, or the Leu286Val Leu392Val mutation. This mutation. the or accomplished by recovering wild type or mutant sequence of RNA using appropriate sources from oligonucleotide primers 989: ggatccggtccacttcgtatgctg, NO:141, 990: ID and SEQ

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ttttttgaattcttaggctatggttgtgttcca, SEQ ID NO:142. This allows cloning of the appropriate mutant or wild type nucleotide sequence corresponding to the hydrophilic acid loop domain at the BamHI and the EcoRI sites within the pGEX-KG vector.

These prokaryotic expression systems allow the holo-protein or various important functional domains of the protein to be recovered as fusion proteins and then used for binding studies, structural studies, functional studies, and for the generation of appropriate antibodies.

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Expression of the ARMP gene in heterologous cell systems can be used to demonstrate structure-function Ligating the ARMP DNA sequence into a relationships. plasmid expression vector to transfect cells is a useful the proteins influence on various method to test Plasmid expression cellular biochemical parameters. vectors containing either the entire, normal or mutant human or mouse ARMP sequence or portions thereof, can be vitro mutagenesis experiments which will inused identify portions of the protein crucial for regulatory function.

The DNA sequence can be manipulated in studies to understand the expression of the gene and its product, to achieve production of large quantities of the protein for functional analysis, for antibody production, and for patient therapy. The changes in the sequence may or may not alter the expression pattern in terms of relative quantities, tissue-specificity and functional properties. Partial or full-length DNA sequences which encode for the ARMP protein, modified or unmodified, may

be ligated to bacterial expression vectors. E. coli can be used using a variety of expression vector systems, e.g., the T7 RNA polymerase/promoter system using two plasmids or by labeling of plasmid-encoded proteins, or by expression by infection with M13 Phage mGPI-2. E . lamba Phage with be used can also coli vectors regulatory sequences, by fusion protein vectors (e.g. lacZ and trpE), by maltose-binding protein fusions, and by glutathione-S-transferase fusion proteins, etc., all of which together with many other prokaryotic expression systems are widely available commercially.

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Alternatively, the ARMP protein can be expressed in insect cells using baculoviral vectors, or in mammalian cells using vaccinia virus or specialized eukaryotic For expression in mammalian cells, expression vectors. heterlogous be ligated to may sequence CDNA promoters, such as the simian varus (SV40) promoter in the pSV2 vector and other similar vectors and introduced into cultured eukaryotic cells, such as COS cells to The stable achieve transient or long-term expression. integration of the chimeric gene construct maintained in mammalian cells by biochemical selection, such as neomycin and mycophoenolic acid.

altered be sequence can DNA The ARMP procedures such as restriction enzyme digestion, fill-in 25 with DNA polymerase, deletion by exonuclease, extension by terminal deoxynucleotide transferase, ligation of synthetic or cloned DNA sequences and site-directed of specific the use with seguence alteration oligonucleotides together with PCR. 30

The cDNA sequence or portions thereof, or a mini gene consisting of a cDNA with an introl and its own is introduced into eukaryotic expression promoter, vectors by conventional techniques. These the cDNA in eukaryotic permit the transcription of cells by providing regulatory sequences that initiate and enhance the transcription of the cDNA and polyadenylation. The splicing and its proper used. also be promoter can endogenous ARMP gene have different within vectors Different promoters activities which alters the level of expression of the In addition, certain promoters can also modulate function such as the glucocorticoid-responsive promoter from the mouse mammary tumor virus.

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the vectors listed contain selectable Some of markers or neo bacterial genes that permit isolation of cells by chemical selection. Stable long-term vectors episomal, cells be maintained in as can entities using regulatory elements replicating viruses. Cell lines can also be produced which have integrated the vector into the genomic DNA. manner, the gene product is produced on a continuous basis.

Vectors are introduced into recipient cells by
various methods including calcium phosphate, strontium
phosphate, electroporation, lipofection, DEAE dextran,
microinjection, or by protoplast fusion. Alternatively,
the cDNA can be introduced by infection using viral
vectors.

30 Using the techniques mentioned, the expression vectors containing the ARMP gene or portions thereof can

be introduced into a variety of mammalian cells from other species or into non-mammalian cells.

The recombinant cloning vector, according to this invention, comprises the selected DNA of the sequences of this invention for expression in a suitable The DNA is operatively linked in the vector to an sequence in the recombinant expression control molecule so that normal and mutant ARMP protein can be The expression control sequence may selected from the group consisting of sequences that control the expression of genes of prokaryotic or eukaryotic cells and their viruses and combinations sequence The expression control thereof. selected from the group consisting of the lac system, the trp system, the tac system, the trc system, major operator and promoter regions of phage lambda, the control region of the fd coat protein, early and late promoters of SV40, promoters derived from polyoma, adenovirus, retrovirus, baculovirus, simian virus, phosphoglycerate kinase promoter, yeast acid phosphatase 20 promoters, yeast alpha-mating factors and combinations thereof.

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The host cell which may be transfected with the vector of this invention may be selected from the group consisting of E. coli, pseudomonas, bacillus subtillus, bacillus stearothermophilus, or other bacili; bacteria, yeast, fungi, insect, mouse or other animal, plant hosts, or human tissue cells.

For the mutant ARMP DNA sequence similar systems 30 are employed to express and produce the mutant protein. Antibodies to Detect ARMP

Antibodies to epitopes with the ARMP protein can be raised to provide information on the characteristics of Generations of antibodies would enable the proteins. the visualizations of the proteins in cells and tissues using Western blotting. In this technique, proteins are run on polyacrylamide gel and then transferred onto are then These membranes nitrocellulose membranes. incubated in the presence of the antibody (primary), then following washing are incubated to a secondary antibody which is used for detection of the proteinprimary antibody complex. Following repeated washing, the entire complex is visualized using colourimetric or chemiluminescent methods.

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Antibodies to the ARMP protein also allow for the use of immunocytochemistry and immunofluorescence techniques in which the proteins can be visualized directly in cells and tissues. This is most helpful in order to establish the subcellular location of the protein and the tissue specificity of the protein.

In order to prepare polyclonal antibodies, fusion 20 proteins containing defined portions or all of the ARMP protein can be synthesized in bacteria by expression of in a suitable cloning corresponding DNA sequences The protein can then be purified, coupled to a vehicle. with Freund's adjuvant carrier protein and mixed 25 the antigenic response by help stimulate rabbits) and injected into rabbits or other laboratory Alternatively, protein isolated can be animals. expressing protein. cells the from cultured bi-weekly injections at Following booster 30 intervals, the rabbits or other laboratory animals

The and the sera isolated. then bled are be used directly or purified prior to use, can affinity including various methods by chromatography, Protein A-Sepharose, Antigen Sepharose, The sera can then be used to Anti-mouse-Ig-Sepharose. probe protein extracts run on a polyacrylamide gel to identify the ARMP protein. Alternatively, synthetic peptides can be made to the antigenic portions of the protein and used to inoculate the animals.

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antibodies. cells produce monoclonal ARMP actively expressing the protein are cultured or isolated from tissues and the cell membranes isolated. The membranes, extracts or recombinant protein extracts, containing the ARMP protein, are injected in Freund's adjuvant into mice. After being injected 9 times over a three week period, the mice spleens are removed and resuspended in a phosphate buffered saline (PBS). The spleen cells serve as a source of lymphocytes, some of are producing antibody of the appropriate which These are then fused with a permanently specificity. growing myeloma partner cell, and the products of the fusion are plated into a number of tissue culture wells in the presence of a selective agent such as HAT. wells are then screened to identify those containing cells making useful antibody by ELISA. These are then freshly plated. After a period of growth, these wells are again screened to identify antibody-producing cells. Several cloning procedures are carried out until over the wells contain single clones which are 90% of positive for antibody production. From this procedure a 30 stable line of clones is established which produce the antibody. The monoclonal antibody can then be purified by affinity chromatography using Protein A Sepharose, ion-exchange chromatography, as well as variations and combinations of these techniques.

In situ hybridization is another method used to situ Inthe expression of ARMP protein. detect hybridization hybridization relies upon the specifically labeled nucleic acid probe to the cellular RNA in individual cells or tissues. Therefore, allows the identification of mRNA within intact tissues, In this method, oligonucleotides such as the brain. corresponding to unique portions of the ARMP gene are used to detect specific mRNA species in the brain.

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is anesthetized and a rat method this In transcardially perfused with cold PBS, followed by perfusion with a formaldehyde solution. The brain or other tissues is then removed, frozen in liquid and cut into thin micron sections. nitrogen, slides and incubated in placed on sections are Following rinsing in DEP, water and proteinase K. 20 in prehybridization slides are placed ethanol, the buffer. A radioactive probe corresponding to the primer is made by nick translation and incubated with the incubation and air sectioned brain tissue. After visualized by areas are the labeled drying, on the tissue sample spots autoradiography. Dark indicate hybridization of the probe with brain mRNA which demonstrates the expression of the protein.

Antibodies may also be used coupled to compounds and/or therapeutic uses diagnostic 3.0 radionuclides for imaging and therapy and liposomes for the targeting of compounds to a specific tissue location.

Isolation and Purification of E5-1 protein

The E5-1 protein may be isolated and purified by the types of methods described above for the ARMP protein.

The protein may also be prepared by expression of the E5-1 cDNA described herein in a suitable host. protein is a preferably expressed as a fusion protein by encoding cDNA sequence to its ligating containing the coding sequence for another suitable The fusion protein is expressed and peptide, e.g., GST. recovered from prokaryotic cells such as bacterial or baculovirus cells or from eukaryotic cells. Antibodies to ARMP, by virtue of portions of amino acid sequence identity with E5-1, can be used to purify, attract and bind to E5-1 protein and vice versa.

Transgenic Mouse Model of E5-1 Related Alzheimer's Disease

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An animal model of Alzheimer's Disease related to mutations of the E5-1 gene may be created by methods analogous to those described above for the ARMP gene.
Antibodies

its structural similarity with the 25 to used for the E5-1 protein may be ARMP. the antibodies probes, peptides, or development of various peptides within it which may recognize both the E5-1 and the ARMP gene and gene products, respectively. As a protein homologue for the ARMP, the E5-1 protein 30

may be used as a replacement for a defective ARMP gene

product. It may also be used to elucidate functions of the ARMP gene in tissue culture and vice versa.

Screening for Alzheimer's
Disease Linked to Chromosome 1

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Screening for Alzheimer's Disease linked to mutations of the E5-1 gene may now be conveniently carried out.

General screening methods are described above in relation to the described mutations in the ARMP gene. These described methods can be readily applied and adapted to detection of the described chromosome 1 mutations, as will be readily understood by those skilled in the art.

In accordance with one embodiment of the invention, the Asn141Ile mutation is screened for by PCR amplification of the surrounding DNA fragment using the primers:

1041: 5'-cattcactgaggacacacc (end-labelled) SEQ ID 20 NO:163 and

1042: 5'-tgtagagcaccaccaaga (unlabelled) SEQ ID NO:164.

Any tissue with nucleated cell may be examined. The amplified products are separated by electrophoresis and an autoradiogram of the gel is prepared and examined for mutant bands.

In accordance with a further embodiment, the Met239Val mutation is screened for by PCR amplification of its surrounding DNA fragment using the primers:

1034: 5'-gcatggtgtgcatccact SEQ ID NO:165 and

1035: 5'-ggaccactctgggaggta SEQ ID NO:166.

The amplified products are separated and an autoradiogram prepared as described above to detect mutant bands.

The same primer sets may be used to detect the mutations by means of other methods such as SSCP, chemical cleavage, DGGE, nucleotide sequencing, ligation chain reaction and allele specific oligonucleotides. As will be understood by those skilled in the art, other suitable primer pairs may be devised and used.

primary event, the as inherited cases, 10 a secondary event due cases as in non-inherited and to the disease state, abnormal processing of E5-1, ARMP, APP or proteins reacting with E5-1, APP or ARMP, may abnormal detected as be can This occur. phosphorylation, glycoslyation, glycation amidation or 15 proteolytic cleavage products in body tissues or fluids, e.g., CSF or blood.

Therapies

important aspect of the biochemical studies using the genetic information of this invention is the 20 development of therapies to circumvent or overcome the ARMP gene defect, and thus prevent, treat, control serious symptoms or cure the disease. In view of expression of the ARMP gene in a variety of tissues, one has to recognize that Alzheimer's Disease may not be 25 restricted to the brain. Alzheimer's Disease manifests itself as a neurological disorder which in one of its forms is caused by a mutation in the ARMP gene, but such manifest may be caused by mutations in other organ such as the liver, releasing factors which 30 tissues, activity and ultimately cause brain affect the

Alzheimer's Disease. Hence, in considering various therapies, it is understood that such therapies may be targeted at tissue other than the brain, such as heart, placenta, lung, liver, skeletal muscle, kidney and pancreas, where ARMP is also expressed.

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The effect of these mutations in E5-1 and ARMP is a gain of novel function which causes aberrant processing Amyloid Precursor Protein into $A\beta$ peptide, (APP) homeostasis, phosphorylation and abnormal Therapy to reverse this will be small apoptosis. molecules (drugs) recombinant proteins, etc. which block the aberrant function by altering the structure of the mutant proteins, etc. which block the aberrant function the mutant protein, altering the structure of enhancing its metabolic clearance or inhibiting binding mutant protein, enhancing of ligands to the metabolic clearance or inhibiting binding of ligands to the mutant protein, or inhibiting the channel function of the mutant protein. The same effect might be gained by inserting a second mutant protein by gene therapy similar to the correction of the "Deg 1(d)" and "Mec 4(d)" mutations in C. elegans by insertion of mutant transgenes. Alternatively over expression of wild type E5-1 protein or wild type ARMP or both may correct the defect. This could be the administration of drugs or proteins to induce the transcription and translation the catabolism of the native or inhibit E5-1 and ARMP proteins. It could also be accomplished by infusion of recombinant proteins or by gene therapy with vectors causing expression of the normal protein at a high level.

Rationale for Therapeutic, Diagnostic, and Investigational Applications of the ARMP Gene and Gene Products as They Relate to the Amyloid Precursor Protein

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The Aeta peptide derivatives of APP are neurotoxic (Selkoe et al, 1994). APP is metabolized by passages through the Golgi network and then to secretory pathways via clathrin-coated vesicles with subsequent passage to the plasma membrane where the mature APP is cleaved by lpha-secretase to a soluble fraction (Protease Nexin II) plus a non-amyloidogenic C-terminal peptide (Selkoe et Alternatively, mature al. 1995, Gandy et al., 1993). APP can be directed to the endosome-lysosome pathway where it undergoes beta and gamma secretase cleavage to produce the Aeta peptides. The phosphorylation state of the cell determines the relative balance of lpha-secretase (amyloidogenic pathways (non-amyloidogenic) Αβ or The phosphorylation (Gandy et al., 1993). pathway) state of the cell can be modified pharmacologicially by phorbol esters, muscarinic agonists and other agents, mediated by cytosolic factors appears to be and (especially protein kinase C) acting upon an integral membrane protein in the Golgi network, which we propose to the ARMP, and members of the homologous family (all phosphorylation consensus several carry which of sequences for protein kinease C). Mutations in the ARMP structure cause alterations in the gene will function of the ARMP gene product leading to defective interactions with regulatory elements (e.g., kinase C) or with APP, thereby promoting APP to be directed to the amyloidogenic endosome-lysosome pathway.

Environmental factors (viruses, toxins, and aging, etc.) may also have similar effects on ARMP. Alzheimer's Disease, the phosphorylation state of ARMP can be altered by chemical and biochemical agents (e.g. compounds) which other drugs, peptides and the activity of protein kinase C and other protein the activity of protein which alter kinase, or modify the availability of which phosphatases, or postranslationally modified. The be ARMP to interactions between kinases and phosphatases with gene products (and the products of the ARMP homologues), and the interactions of the ARMP products with other proteins involved in the trafficking of the APP within the Golgi network can be modulated to decrease trafficking of Golgi vesicles to the endosomepathway thereby promoting Αβ peptide lvsosome Such compounds will include: peptide production. analogues of APP, ARMP, and homologues of ARMP as well other interacting proteins, lipids, sugars, agents which promote differential glycosylation of ARMP and its homologues; agents which alter the biologic ARMP half-life of messenger RNA or protein of including antibodies and antisense homologues oligonucleotides; and agents which act upon ARMP 25 transcription.

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The effect of these agents in cell lines and whole animals can be monitored by monitoring: transcription; modification of ARMP translation; post-translational glycoslyation); and phosphorylation orintracellular trafficking of ARMP and its homologues and extracellular through various intracellular

compartments. Methods for these studies include Western and Northern blots; immunoprecipitation after metabolic labeling (pulse-chase) with radio-labeled methionine and The effect of these and imminohistochemistry. agents can also be monitored using studies which examine the relative binding affinities and relative amounts of ARMP gene products in interactions with protein kinease C and/or APP using either standard binding affinity Western blots using assays or co-precipitation and antibodies to protein kinease C, APP or ARMP and its The effect of these agents can also be homologues. monitored by assessing the production of $A\beta$ peptides by exposure to the putative after before and therapeutic agent (Huang et al., 1993). The effect can also be monitored by assessing the viability of cell lines after exposure to aluminum salts and to Aβ neurotoxic in through to be are peptides which Finally, the effect of these Alzheimer's Disease. agents can be monitored by assessing the cognitive function of animals bearing: their normal genotype at APP or ARMP homologues; bearing human APP transgenes (with or without mutations); or bearing human ARMP transgenes (with or without mutations); or a combination of all of these.

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Rationale for Therapeutic, Diagnostic, and Investigational Applications of the ARMP Gene, the E5-1 Gene and their Products

The ARMP gene product and the E5-1 gene product have amino acid sequence homology to human ion channel proteins and receptors. For instance, the E5-1 protein shows substantial homology to the human sodium channel

 α -subunit (E=0.18, P=0.16, identities=22 - 27% over two regions of at least 35 amino acid residues) using the BLASTP paradigm of Atschul et al. 1990. Other diseases hyperkalemic hyperthermia and maliqnant (such periodic paralysis in humans and the neurodegenerative of mechanosensory neurons in C. elegans) arise through channels or receptor proteins. mutations ion Mutation of the ARMP gene or the E5-1 gene could affect similar functions and lead to Alzheimer's Disease and other psychiatric and neurological diseases. Based upon this, a test for Alzheimer's Disease can be produced to detect an abnormal receptor or an abnormal ion channel function related to abnormalities that are acquired or inherited in the ARMP gene and its product or in one of the homologous genes such as E5-1 and their products. This test can be accomplished either in vivo or in vitro and/or channel fluxes of ion measurements by transmembrane voltage or current fluxes using patch clamp, voltage clamp and fluorescent dyes sensitive to transmembrane voltage. intracellular calcium or Defective ion channel or receptor function can also be of second assayed by measurements of activation cGMP tyrosine cyclic AMP, messengers such as intracellular Ca2+ kinases, phosphates, increases in etc. Recombinantly made proteins may levels, be reconstrued in artificial membrane systems to study Therapies which affect ion channel conductance. Alzheimer's Disease (due to acquired/inherited defects in the ARMP gene or E5-1 gene; due to defects in this disease such leading to pathways other APP; and due to environmental mutations in

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by analysis of be tested agents) can ion channel abnormal modify an ability to receptor function induced by mutation in the ARMP gene or in one of its homologues. Therapies could also be tested by their ability to modify the normal function of receptor capacity of the channel or gene products and its homologues. Such assays can cells expressing cultured performed on mutant ARMP genes/gene products endogenous normal or Such studies can be E5-1 genes/gene products. performed in addition on cells transfected with vectors capable of expressing ARMP, parts of the ARMP gene and gene product, mutant ARMP, E5-1 gene, parts of the E5-1 gene and gene product, mutant E5-1 gene or another homologue in normal or mutant form. Therapies Alzheimer's Disease can be devised to modify an abnormal ion channel or receptor function of the ARMP gene or E5-Such therapies can be conventional drugs, 1 gene. peptides, sugars, or lipids, as well as antibodies or other ligands which affect the properties of the ARMP or E5-1 gene product. Such therapies can also be performed by direct replacement of the ARMP gene and/or E5-1 gene In the case of an ion channel, the by gene therapy. gene therapy could be performed using either mini-genes (cDNA plus a promoter) or genomic constructs bearing genomic DNA sequences for parts or all of the ARMP gene. Mutant ARMP or homologous gene sequence might also be used to counter the effect of the inherited or acquired abnormalities of the ARMP gene as has recently been done for replacement of the mec 4 and deg 1 in C. elegans (Huang and Chalfie, 1994). The therapy might also be

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directed at augmenting the receptor or ion channel function of the homologous genes such as the E5-1 gene, in order that it may potentially take over the functions of the ARMP gene rendered defective by acquired or inherited defects. Therapy using antisense oligonucleotides to block the expression of the mutant ARMP gene or the mutant E5-1 gene, coordinated with gene replacement with normal ARMP or E5-1 gene can also be applied using standard techniques of either gene therapy or protein replacement therapy.

Protein Therapy

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Treatment of Alzheimer's Disease can be performed by replacing the mutant protein with normal protein, or by modulating the function of the mutant protein. Once the biological pathway of the ARMP protein has been completely understood, it may also be possible to modify the pathophysiologic pathway (e.g., a signal transduction pathway) in which the protein participates in order to correct the physiological defect.

To replace the mutant protein with normal protein, or with a protein bearing a deliberate counterbalancing mutation it is necessary to obtain large amounts of pure ARMP protein or E5-1 protein from cultured cell systems which can express the protein. Delivery of the protein to the affected brain areas or other tissues can then be accomplished using appropriate packaging or administrating systems.

Gene Therapy

Gene therapy is another potential therapeutic approach in which normal copies of the ARMP gene are introduced into patients to successfully code for normal

protein in several different affected cell types. The gene must be delivered to those cells in a form in which it can be taken up and code for sufficient protein to provide effective function. Alternatively, in some neurologic mutants it has been possible to prevent disease by introducing another copy of the homologous gene bearing a second mutation in that gene or to alter mutation, or use another gene to block its effect.

Retroviral vectors can be used for somatic cell gene therapy especially because of their high efficiency of infection and stable integration and expression. The targeted cells however must be able to divide and the expression of the levels of normal protein should be high because the disease is a dominant one. The full length ARMP gene can be cloned into a retroviral vector and driven from its endogenous promoter or from the retroviral long terminal repeat or from a promoter specific for the target cell type of interest (such as neurons).

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Other viral vectors which can be used include adeno-associated virus, vaccinia virus, bovine papilloma virus, or a herpesvirus such as Epstein-Barr virus.

Gene transfer could also be achieved using non-viral means requiring infection in vitro. This would include calcium phosphate, DEAE dextran, electroporation, and protoplast fusion. Liposomes may also be potentially beneficial for delivery of DNA into a cell. Although these methods are available, many of these are lower efficiency.

Antisense based strategies can be employed to explore ARMP gene function and as a basis for

therapeutic drug design. The principle is based hypothesis that sequence-specific suppression of expression can be achieved by intracellular gene hybridization between mRNA and a complementary antisense The formation of a hybrid RNA duplex may then processing/transport/translation interfere with the and/or stability of the target ARMP mRNA. Hybridization is required for the antisense effect to occur, however the efficiency of intracellular hybridization is low and therefore the consequences of such an event may not be Antisense strategies may use a variety very successful. antisense the use of including of approaches RNA and antisense oligonucleotides, injection of expression transfection of antisense RNA vectors. Antisense effects can be induced by control sequences, however, the extent of phenotypic changes are effects induced highly variable. Phenotypic by antisense effects are based on changes in criteria such as protein levels, protein activity measurement, target mRNA levels. Multidrug resistance is a useful study molecular events associated model to phenotypic changes due to antisense effects, since the multidrug resistance phenotype can be established by expression of a single gene mdrl (MDR gene) encoding for P-glycoprotein.

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Transplantation of normal genes into the affected area of the patient can also be useful therapy for Alzheimer's Disease. In this procedure, a normal hARMP protein is transferred into a cultivable cell type such as glial cells, either exogenously or endogenously to the patient. These cells are then injected

serotologicially into the disease affected tissue(s). This is a known treatment for Parkinson's disease.

Immunotherapy is also possible for Alzheimer's Antibodies can be raised to a mutant Disease. administered portion thereof) and then protein (or and mutant protein the or block bind expression Simultaneously, effects. deliterious could encouraged. be protein product normal in the form of a one time Administration could be immunogenic preparation or vaccine immunization. 10 as prepared composition may be immunogenic liquid solutions or emulsions. The injectables, as mixed with pharmaceutically be may protein ARMP acceptable excipients compatible with the protein. include water, saline, dextrose, excipients may 15 combinations thereof. The and glycerol, ethanol immunogenic composition and vaccine may further contain auxiliary substances such as emulsifying agents enhance effectiveness. Immunogenic adjuvants to administered vaccines may be and compositions 20 parenterally by injection subcutaneously intramuscularly.

The immunogenic preparations and vaccines are administered in such amount as will be therapeutically effective, protective and immunogenic. Dosage depends on the route of administration and will vary according to the size of the host.

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Similar gene therapy techniques may be employed with respect to the $\it E5-1$ gene.

The above disclosure generally describes the present invention. A more complete understanding can be

specific following reference to the obtained by These examples are described solely for examples. purposes of illustration and are not intended to limit Changes in the form and the scope of the invention. contemplated equivalents are of substitution circumstances may suggest or render expedient. specific terms have been employed herein, such terms are intended in a descriptive sense and not for purposes of limitations.

10 Example 1. Development of the Genetic, Physical "contig" and Transcriptional Map of the Minimal Co-Segregating Region

The CEPH Mega YAC and the RPCI PAC human total for searched libraries were genomic DNA ' containing genomic DNA fragments from the AD3 region of chromosome 14q24 & using oliginucleotide probes for each of the ## SSR marker loci used in the genetic linkage studies as well as \## additional markers depicted in 20 Figure 1a (Albertsen \et al., 1990; Chumakov et al., 1992; Ioannu et al., 1994). The genetic map distances between each marker are depicted above the contig, and are derived from published \data (NIH/CEPH Collaborative Mapping Group, 1992; Wang, 1992; Weissenbach, J. et al., 25 1992 Gyapay, G et al., 1994). Clones recovered for each of the initial marker loci were arranged into an ordered series of partially overlapping clones ("contig") using four independent methods. First, sequences representing the ends of the YAC insert were isolated by inverse PCR 30 (Riley et al., 1990), and hybridized to Southern blot panels containing restriction digests of \DNA from all of the YAC clones bearing overlapping sequences. Second,

inter-Alu \PCR was performed on each YAC, and the resultant band patterns were compared across the pool of recovered YA clones in order to identify other clones bearing over apping sequences (Bellamne-Chartelot al., 1992; Chumakov et al; 1992). Third, to improve the specificity of the Alu-PCR fingerprinting, we restricted RsaI, amplified the YAC DNA with HaeIII or restriction products with both Alu and L1H consensus primers, and resolved the products by polyacrylamide gel as additional STSs were \Finally, electrophoresis. generated during the search for transcribed sequences, these STSs were also used to identify overlaps. The resultant contig was complete except for a single discontinuity between \ YAC932C7 bearing D14S53 YAC746B4 containing D14S61. The physical map order STSs within the contig was largely in the with the \genetic linkage map for this accordance region (NIH/CEPH Collaborative Mapping Group, 1992; Wang, Z., Webber, J.L., 1990; Weissenbach, J. et al., 1992; Gyapay, G. et al., 1994). However, as with the genetic maps, we were unable to unambiguously resolve the loci within the order of the relative D14S43/D14S71 cluster and the D14S76/D14S273 cluster. suggest that D149277 is telomeric clones PAC1 D14S268, whereas genetic maps have \suggested the reverse Furthermore, a few STS probes failed to detect hybridization patterns in at least one YAC clone which, on the basis of the most parsimonious consensus physical map and from the genetic map, would have been predicted to contain that STS. For instance, the D14S268 (AFM265) and RSCAT7 STSs are absent from YAC788H12 (Figure 3).

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Because \these results are reproducible, and occurred with several different STS markers, these results most of small interstitial likely reflect the presence deletions with one of the YAC clones.

Example 2. Cumulative two-point lod scores for hromosome 14q24.3 markers.

Genotypes of each polymorphic microsatellite marker locus were determined by PCR from 1000ng of genomic DNA available affected and unaffected pedigree all members as previously described (St. George-Hyslop, P et using primer sequences specific for each al., 1992; (Weissenbach, et locus J microsatellite The normal population 1994). Gyapay, G al., et frequency of each allele was determined using spouses and other neurologically normal subjects from the same ethnic groups, but did not differ significantly from populations Caucasian established for mixed those (Weissenbach, J. et al., 1992; Gyapay, G. et al., 1994). The maximum likelihood calculations assumed an age of onset correction, marker allele frequencies derived from mixed Caucasian subjects, published of series estimated allele frequence for the AD3 mutation 1:1000 as previously described (St. George-Hyslop, P. et The analyses were repeated using equal al., 1992). phenotype using frequencies, and allele marker information only from affected pedigree members previously described to ensure that inaccuracies in the estimated parameters used in the maximum likelihood calculations did not misdirect the analyses (St. George-30 Hyslop, P. et al., 1992). These supplemental analyses not significantly alter either the

supporting linkage, or the discovery of recombination events.

Example 3. Haplotypes between flanking markers segregated with AD3 in FAD pedigrees

Extended haplotypes between the centromeric and telomeric flanking markers on the parental copy of chromosome 14 segregating with AD3 in fourteen early onset FAD pedigrees (pedigrees NIH2, MGH1, Tor1.1, FAD4, FAD1, MEX1, and FAD2 show pedigree specific lod scores \geq 10 least one marker between D14S258 and +3.00 with at partial haplotypes (boxed) are Identical D14S53). bearing disease regions of the observed in two chromosome segregating in several pedigrees of similar ethnic origin. In region A, shared alleles are seen at 15 D14S268 ("B": allele size = 126 bp, allele frequence in normal Caucasians = 0.04; "C": size = 124 bp, frequency = 0.38); D14S277 ("B": size = 156 bp, frequency = 0.19; "C": size - 154 bp, frequency = 0.33); and RSCAT6 ("D": size = 111bp, frequency 0.25; "E" size = 109 bp, 20 frequency = 0.20; "F" size = 107 bp, frequency = 0.47). In region B, alleles of identical size are observed at D14S43 ("A": size = 193bp, frequency = 0.01; "D": size 187 bp, frequency = 0.12; "E" size = 185bp, frequency = 0.26; "I" size = 160 bp, frequency = 0.38); D14S273 25 ("3": size = 193 bp, frequency = 0.38; "4" size = 191 bp, frequency = 0.16; "5": size = 189 bp, frequency = 0.34; "6": size = 187 bp, frequency = 0.02) and D14S76 ("1": size = bp, frequency = 0.01; "5": size = bp, frequency = 0.38; "6": size = bp, frequency = 0.07, "9": 30 The ethnic origins of size = bp, frequency = 0.38). Ashk = Askenazi each pedigree are abbreviated as:

Jewish; Ital = Southern Italian; Angl = Anglo-Saxon-Celt; FrCan = French Canadian; Jpn = Japanese; Mex = Mexican Caucasian; Ger = German; Am = American Caucasian. The type of mutation detected is depicted by the amino acid substitution and putative condon number or by ND where no mutation has been detected because a comprehensive survey has not been undertaken due to the absence of a source of mRNA for RT-PCR studies.

Example 4. Recovery of transcribed sequences from the AD3 interval.

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sequences encoded in the AD3 Putative transcribed direct using either recovered interval were in which short cDNA fragments hybridization method generated from human brain mRNA were hypridized to immobilized cloned genomic DNA fragments (Rommens, JM et The resultant short putatively transcribed al., 1993). to recover probes sequences were used as transcripts from human brain cDNA libraries (Stratagene, La Jolla). The physical locations of the original short clone and of the subsequently acquired longer cDNA clones were established by analysis of the hybridization pattern generated by hybridizing the probe to Southern blots containing a panel of EcoRI digested total DNA samples isolated from individual YAC clones within the The nucleotide sequence of each of the longer cDNA clones was determined by automated cycle sequencing (Applied Biosystems Inc., CA), and compared to other sequences in nucleotide and protein databases using the blast algorithm (Atschul, SF et al., 1990). numbers for the transcribed sequences in this report are L40391, L40392, L40393, L40394, L40395, L40396, L40397, L40398, L40399, L40400, L40401, L40402, and L40403.

Example 5. Locating mutations in the ARMP gene using restriction enzymes.

5 The presence of Ala 246 Glu mutation which creates a Ddel restriction site was assayed in genomic DNA by (5! end labelled primer 849 using the atctccggcaggcatatct-3') SEQ ID NO:129 and the unlabelled primer 892 (5'-tgaaatcacagccaagatgag-3') SEQ ID NO:130 10 to amplify an 84bp genomic exon fragment using 100ng of genomic DNA template, 2mM MgCl₂, 10 pMoles of each primer, 0.5U Taq polymerase, 250 uM dNTPs for 30 cycles of 95°C x 20 seconds, 60°C X 20 seconds, 72°C X 5 seconds. The products were incubated with an excess of 15 hours manufacturers according to the for 2 DdeI protocol, and the resulting restriction fragments were resolved on a 6% nondenaturing polyacrylamide gel and visualized by autoradiography. The presence of the mutation was inferred from the clevage of the 84bp 20 fragment due to the presence of a DdeI restriction site. affected members of the FAD1 pedigree (filled All symbols) and several at-risk members ("R") carried the None of the obligate escapees (those DdeI site. individuals who do not get the disease, age > 70 years), 25 and none of the normal controls carried the DdeI mutation.

Example 6. Location mutation in the ARMP gene using allelle specific oligonucleotides.

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The presence of the Cys 410 Tyr mutation was assayed using allele specific oligonucleotides. 100ng of genomic DNA was amplified with the exonic sequence

primer 885 (5'-tggagactggaacacaac-3') SEQ ID NO:127 and (5'intronic sequence primer 893 opposing gtgtggccagggtagagaact-3') SEQ ID NO:128 using the above reaction conditions except 2.5 mM MgCl₂, conditions of 94°C X 20 seconds, 58°C X 20 seconds, and The resultant 216bp genomic 72°C for 10 seconds). fragment was denatured by 10-fold dilution in 0.4M NaOH, and was vacuum slot-blotted to duplicate 25mM EDTA, The end-labelled "wild-type" primer nylon membranes. 890 (5'-ccatagcctgtttcgtagc-3') SEQ ID NO:131 and the (5'-891 primer "mutant" end-labelled ccatagcctAtttcgtagc-3') SEQ ID NO:132 were hybridized to separate copies of the slot-blot filters in 5 X SSC, 5X Denhardt's, 0.5% SDS for 1 hour at 48°C, and then washed successively in 2 X SSC at 23°C and 2 X SSC, 0.1% SDS at All testable 50°C and then exposed to X-ray film. affected members as well as some at-risk members of the AD3 (shown) and NIH2 pedigrees (not shown) possessed the Cys 410 Tyr mutation. Attempts to detect the Cys 410 Try mutation by SSCP revealed that a common intronic sequence polymorphism migrated with the same SSCP pattern.

Example 7. Northern hybridization demonstrating the expression of ARMP protein mRNA in a variety of tissues.

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isolated from cytoplasmic RNA was Total (including heart, brain, and various tissue samples liver, placenta, lung, regions of different kidney and pancreas) obtained from skeletal muscle, surgical pathology using standard procedures such as was then electrophoresed CsCl purification. The RNA to permit size fractionation. a formaldehyde gel on

was nitrocellulose membrane prepared and the The 32 p membrane. then transferred onto the RNA was and added to were prepared probes labelled CDNA hybridization membrane in order for the occur. After washing, RNA to the probe the and placed plastic film wrapped in membrane was imaging cassettes containing The X-ray film. autoradiographs were then allowed to develop for one to The positions of the 28S and 18S rRNA several days. Sizing was established by bands indicated. are comparison to standard RNA markers. Analysis of the autoradiographs revealed a prominent band at 3.0kb in These northern blots demonstrated the ARMP gene size. is expressed in all of the tissues examined.

15 Example 8: Eukaryotic and Prokaryotic Expression Vector Systems

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Eukaryotic and prokaryotic expression systems have been generated using two different classes of nucleotide cDNA sequence inserts. In the first class, termed full-length constructs, the entire ARMP sequence was inserted into the expression plasmid in the correct orientation, and included both the natural 5' UTR and 3' UTR sequences as well as the entire open reading frames The open reading frame. nucleotide sequence cassette which allows either wild type open reading frame to be included in the expression system or alternatively, single combination of double mutations can be inserted into the open reading frame. This was accomplished by removing a restriction fragment from the wild type open reading frame using the enzymes NarI and PflmI and replacing it

generated by fragment similar with а nucleotide bears the which and transcriptase PCR sequence encoding either the Met146Leu mutation or the A second restriction fragment was Hys163Arg mutation. removed from the wild type normal nucleotide sequence for the open reading frame by cleavage with the enzymes PflmI and NcoI and replaced with restriction fragments bearing either the nucleotide sequence encoding the Ala246Glu mutation, or the Ala260Val mutation or the Ala285Val mutation or the Leu286Val mutation, or the mutation. Tyr Leu392Val mutation, or the Cys410 Finally, a third variant bearing combinations of either the Met146Leu or His163Arg mutations in tandem with the remaining mutations by linking the Narl-PflmI fragment bearing these mutations and the PflmI-NcoI fragment bearing the remaining mutations.

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A second variant of cDNA inserts bearing cDNA sequences was constructed by mutant or type removing from the full-length cDNA the 5' UTR and part The 5' UTR sequence was of the 3' UTR sequences. replaced with a synthetic oligonucleotide containing a restriction site and a Kozak initiation site KpnI ggtaccgccaccatgacagaggtacctgcac) (oligonucleotide 969: 3' UTR was replaced with an The ID NO:139. SEQ oligonucleotide corresponding to position 2566 of the cDNA and bears an artificial EcoRI site (oligonucleotide 970:gaattcactggctgtagaaaaagac) SEQ ID NO:140. variants of this construct were then made by inserting the same mutant sequences described above at the NarI-PflmI fragment, and at the PsImI-NcoI sites described above.

For eukaryotic expressions, these various constructs bearing wild type and mutant sequences were cloned into the expression vector pZeoSV (invitrogen). For prokaryotic expression, two constructs were using the gluthathione S-transferase fusion vector pGEX-The inserts which have been attached to the GST are the nucleotide nucleotide sequence same generated with the described above sequence oligonucleotide primers 969, SEQ ID NO:139 and 970, SEQ ID NO:140, bearing either the normal open reading frame nucleotide sequence or bearing a combination of single and double mutations as described above. This construct allows expression of the full-length protein in mutant and wild type variants in prokaryotic cell systems as a GST fusion protein which will allow purification of the full-length protein followed by removal of the fusion product by thrombin digestion. The second prokaryotic cDNA construct was generated to create a fusion protein with the same vector, and allows the production of the amino acid sequence corresponding to the hydrophillic acid loop domains between TM6 and TM7 the full-length protein, as either a wild type sequence (thus a wild type amino acid nucleotide sequence for fusion proteins) or as а mutant sequence bearing either the Ala285Val mutation, or Leu286Val mutation, Leu392Val mutation. the or the This accomplished by recovering wild type or was mutant sequence from appropriate sources of RNA using the oligonucleotide primers 989:ggatccggtccacttcgtatgctg SEQ ID NO:141, and 990:tttttttgaattcttaggctatggttgtgttcca SEQ ID NO:142. This allows cloning of the appropriate

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mutant or wild type nucleotide sequence corresponding to the hydrophillic acid loop domain at the BamHI and the EcoRI sites within the pGEX-KG vector.

These prokaryotic expression systems allow the holo-protein or various important functional domains of the protein to be recovered as fusion proteins and then used for binding studies, structural studies, functional studies, and for the generation of appropriate antibodies.

10 Example 9: Identification of Three New Mutations in the ARMP Gene.

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Three novel mutations have been identified subjects affected with early onset Alzheimer's Disease. All of these mutations co-segregate with the disease, 15 and are absent from at least 200 normal chromosomes. The three mutations are as follows: a substitution of C by T at position 1027 which results in the substitution of alanine 260 for valine; substitution of C by T at position 1102, which results in the substitution of 20 alanine at 285 by valine; and substitution of C by G at position 1422 which results in the substitution of leucine 392 by valine. Significantly, all of these mutations occur within the acidic hydrophillic loop between putative TM6 and TM7. Two of the mutations 25 (A260V; A285V) and the L286V mutation are also located in the alternative spliced domain.

The three new mutations, like the other mutations, can be assayed by a variety of strategies (direct nucleotide sequencing, Allele specific oligos, ligation polymerase chain reaction, SSCP, RFLPs) using RT-PCR products representing the mature mRNA/cDNA sequence or

We have chosen allele specific oligos. genomic DNA. For the A260V and the A285V mutations, genomic DNA carrying the exon can be amplified using the same PCR PCR methods for the mutation. L286V and primers slot blotted denatured and then products were duplicate nylon membranes using the slot blot protocol described for the C410T mutation.

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The Ala260Val mutation was scored by these blots by hybridization with end-labeled allele-specific oligonucleotides corresponding to the wild type sequence the ID NO:143 or SEO (994:gattagtggttgttttgtg) sequence (995:gattagtggctgttttgtg) SEQ ID mutant by hybridization at 48°C followed by a NO:144 SSC buffer containing 0.1% SDS. The 3 X was scored on these slot blots as mutation Ala285Val described above but using instead the allele-specific type sequence the wild oligonucleotides for (1003:tttttccagctctcattta) SEQ ID NO:145 or the mutant primer (1004:tttttccagttctcattta) SEQ ID NO:146 at 48°C followed by washing at 52°C as above except that the wash solution was 2X SSC.

The Leu392Val mutation was scored by amplification using primers 996 DNA from genomic of the exon 893 ID NO:167 and (aaacttggattgggagat) SEQ (gtgtggccagggtagagaact) SEQ ID NO:128 using standard PCR magnesium the conditions excepting that buffer concentration was 2mM and cycle conditions were 94°C time 10 seconds, 56°C times 20 seconds, and 72°C for 10 The result 200 based pair genomic fragment was seconds. denatured as described for the Cys410Tyr mutation and slot-blotted in duplicate to nylon membranes.

presence or absence of the mutation was then scored by differential hybridization to either a wild type end-labelled oligonucleotide (999:tacagtgttctggttggta) SEQ ID NO:148 or with an end-labeled mutant primer (100:tacagtgttgtggttggta) SEQ ID NO:149 by hybridization at 45°C and then successive washing in 2X SSC at 23° and then at 68°C.

Example 10: Polyclonal Antibody Production

Peptide antigens were synthesized by solid-phase techniques and purified by reverse phase high pressure 10 liquid chromatography. Peptides were covalently linked via disulfide limpet hematoxylin (KLH) to keyhole linkages that were made possible by the addition of a cystein residue at the peptide C-terminus. additional residue does not appear normally 15 protein sequence and was included only to facilitate A total of three rabbits linkage to the KLH molecule. immunized with peptide-KLH complexes for peptide antigen and were then subsequently give booster Antisera were seven day intervals. 20 injections at pooled and IqG for each peptide and collected precipitated with ammonium sulfate. Antibodies were then affinity purified with Sulfo-link agarose (Pierce) coupled with the appropriate peptide. This final non-specific purification is required to remove 25 interactions of other antibodies present in either the pre- or post-immune serum.

The specific sequences to which we have raised antibodies are:

B, A

³⁰ Polyclonal antibody 1: NDNRERQEHNDRRSL (C)-residues 30-45 SEQ ID NO:168

SUB18 SUBIC

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Polyclonal antibody 2: KDGQLIYTPFTEDTE (C)-residues 109-120

SEQ ID NO:169

Polyclonal antibody 3: EAQRRVSKNSKYNAE (C)-residues 304-319

SEQ ID NO:170

Polyclonal antibody 4: SHLGPHRSTPESRAA (C)-residues 346-360 SEQ ID NO:171

The non-native cysteine residue is indicated at the C-These sequences are contained within terminal by (C). various predicted domains of the protein. For example, in potentially and 4 are located 1,3, antibodies functional domains that are exposed to the aqueous media involved in binding to other proteins and may be critical for the development of the disease phenotype. linking short corresponds to a 2 Antibody second situated between the predicted first and transmembrane helices.

Example 11: Identification of two mutations in E5-1 gene.

RT-PCR products corresponding to the E5-1 ORF were 20 generated from RNA of lymphoblasts or frozen post-mortem brain tissue using oligonucleotide primer pairs 1021:5'and SEQ ID NO:172 cagaggatggagaatac ggctccccaaaactgtcat SEQ ID NO:173 (product = 888 bp); and 1071:5'-gccctagtgttcatcaagta ID NO:174 SEQ 25 1022:5'-aaagcgggagccaaagtc SEQ ID NO:175 (product = 826 bp) by PCR using 250 µMol dNTPs, 2.5 mM MgCl2, 10pMol oligunucleotides in 10 μl cycled for 40 cycles of 94°C X 20 seconds, 58°C X 20 seconds, 72°C X 45 seconds. The automated cycle products sequenced by 30 PCR were sequencing (ABI, Foster City, A) and the fluorescent chromatograms were scanned for heterozygous nucleotide substitutions by direct inspection and by the Factura (ver 1.2.0) and Sequence Navigator (ver 1.0.1b15) software packages (data not shown).

Asn141Ile: the A \rightarrow T substitution at nucleotide Bc1I restriction site. The 787 creates a mutation was amplified from 100 ng bearing this 5 genomic DNA using 10pMol of oligonucleotides SEQ ID NO:163 1041: 5'-cattcactaggacacacc 1042: 5'-tgtagagcaccaccaaga labelled) and NO:164 (unlabelled), and PCR reaction conditions below for those described 10 similar to Met239Val. 2µl of the PCR product was restricted Bc1I (NEBL, Beverly, MA) in 10 µl reaction according to the manufacturers' protocol, volume and the products were resolved by non - denaturing polyacrylamide gel electrophoresis. In subjects with 15 wild type sequences, the 114 bp PCR product is cleaved into 68 bp and 46 bp fragments. Mutant sequences cause the product to be cleaved into 53 bp, 46 bp and 15 bp.

Met239Val: The A→G substitution at nucleotide 1080 deletes a NlaIII restriction site, allowing the presence 20 Met239Val mutation to be detected the by amplification from 100 ng of genomic DNA using PCR (10 pMol oligonucleotides 1034:5'-gcatggtgtgcatccact SEQ ID NO:165, 1035:5'-ggaccactctgggaggta SEQ ID NO:166; 0.5 U Tag polymerase, 250μM dNTPS, 1μCi alpha ³²P-dCTP, 1.5 mM 25 MqCI₂, 10µl volume; 30 cycles of 94°C X30 seconds, 58°C X 20 seconds, 72°C X 20 seconds) to generate a 110 bp product. 2ul of the PCR reaction were diluted to 10ul and restricted with 3 U of NlaIII (NEBL, Beverly MA) for 3 hours. The restriction products were resolved by non-30 denaturing polyacrylamide gel electrophoresis

visualized by autoradiography. Normal subjects show cleavage products of 55, 35, 15 and 6bp, whereas the mutant sequence gives fragments of 55, 50 and 6 bp.

Although preferred embodiments of the invention have been described herein in detail, it will be understood by those skilled in the art that variations may be made thereto without departing from the spirit of the invention or the scope of the appended claims.

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	-	RECOMBI	NATION F	RACTION	(0)		<u>.</u>
LOCUS	0.00	0.05	0.10	0.15	0.20	0.30	0.40
D14S63	-00	1.54	3.90	4.38	4.13	2.71	1.08
D14S258	-∞	21.60	19.64	17.19	14.50	8.97	3.81
D14S77		15.18	15.53	14.35	12.50	7.82	2.92
D14S71	-∞	15.63	14.14	12.19	10.10	5.98	2.39
D14S43	-∞	19.36	17.51	15.27	12.84	7.80	3.11
D14S273		12.30	11.52	10.12	8.48	5.04	1.91
D14S61	-∞	26.90	24.92	22.14	18.98	12.05	5.07
D14S53	-∞	11.52	11.41	10.39	8.99	5.73	2.51
D14S48	-∞	0.50	1.05	1.14	1.04	0.60	0.18

TABLE 1

						PEDIGI	PEDIGREE ID							
rocns	N1B2	FaD3	TURI.1	FaD4	RB	FaD1	BIG12	BOW	СООК	603	Tor42	QUE	MEXI	FAD2
D14S83	-	4	7	4		5						-1.	6	2
D14S258	9	9	8	7	4	5	5	9	9		7	9	7	9
D14S268	ນ	ပ	æ	В	ပ	ပ	၁	ပ	ပ	C	C C	B	ပ	
D14S277	၁	ပ	၁	၁	ပ	၁	၁	၁	ပ	∢	⋖	C	8	
D14S786	۵	Q	Э	Е	Œ	ធា	ш	D/F	ш	ш	ப	ш	Ľ.	Q
• D14S77	>	>	×	S		Ь	Ь	×	=))	n	뜨	«
D14578	7	7	-	5	7	7		9	7		: &	7	2	9
6 D14S43	∢	<	 - 	: 		<u>ш</u>	Ω	-	Head		၁	-	Q	၁
D14S273	9	9	е 	5	۰ د	4	4	4	9		9	9	8	3
D14S76	S	S	5	5	\$	9	6	6			6	-	2	2
D14S61	ш	ш	ŋ	Ľ.		_					Ω		1	Ľ
D14S53	Ľ	Œ.	၁	ĹŢ.	Ľ	-	C	Œ,	ш		-	Q	ű.	Ľ
ETHNIC	Ashk	Ashk	Ital	Ital	Ital	Angl	Angl	Angl	Angl	Amer	FrCan	FrCan	Mex	ڻ
MUTATION	C410Y	C410Y C410Y	M146L	M146L	Q	A246E	Q	Q	Q	H163R	H163R	Q	QN	L286V

TABLE 2

	•							Similarities
	No. 1	arget File	_	Kay	Target	Overlap	Match	Percencego
	1 E	armo.con/lo	ng[Frame 1]	1	1	467	465	99.57
	1	10	20	30	40	50	60	70
Human	N- MTE	LPAPLSYPCNU	THE THE EM	RSCNONRER	OZHNDRRSI			
	***	*********	••••••	******	*******	*******	*****	******
Mouse	N- MTE	IPAPLEYFONA	Maedahaay.	ergonde of the conference of t	QQQHDRQRI	DNPEPISNO	POSESSES	VECDREED
	1	10	20	30	4,0	50	60	סל
	71	8,0	90	100	110	120	130	140
	EEL	TLKYGNGTVIDG	LEVEVELCHUV	TEVERLITAVI	YTRIOGOLI	YTPFTEDTE	TVGQRALHS	INVADAL
	***	*********	*********	******	•••••	********	*******	******
		WINCE THE PROPERTY OF THE PROP					PVGORALIES	
	71	80	90	100	110	120	130	140
	141	150	160	170	180	190	200	210
	BVI	VVMTILLVVLYI	KYRCYKVIHAWI	TILLLERII.	Frakialce	VFKTYNVAVI	YTTVALLI	MILGVVOM
	***	*********				*******	*******	*****
		VIKITLLVVLYI						
•	141	150	160	170	180	190	200	210
	211	220	230	240	250	260	270	280
	ISI	HWKGPLRLQQAY	CLINISALMALI	FIXTLPEAT.	nliiavis	VYDLVAVLCI	PROPLEMEN	ETACERNE
	***	**********			******	******	*******	******
•	1Ai	HWKCPLRLOOAY						· -
	211	220	230	240	250	260	270	280
	281	290	300	310	320	330	340	350
	TLF	Paliyss Th iwi	VNMAECOPEA	(YXBOXEVRX	n ae st e res	ODIVEDOO	Corsed (Ca	ORDSHLOP
	74.	******			•••••	*****		******
	TLF.	INVMTEEYILLG						
	281	290	300	310	320	330	340	350
	351	360	370	380	390	400	410	420
	HR.S'	TPESRAAVQELS	issilaced per	RGVXLALAD	Pifybvlvg	Kerkerk	nttiaciv	ATLIGICL
	***		**********	*******	•••••	******	******	*******
		TPESRAAVQELS						
	351	360	370	380	390	400	410	420
	421	430	440	450	460			
	TLL	Liaipkkalpai	PISITECLVY	PATDYLVOP	PHOCEATHO	MI -C	SEQ ID N	O: 2
	XT.I.	LLATYKKGXPAX	PIRITEGEVEY	PATTYLVA	MOOTA FED	TY7 -C	SEQ ID N	iO: 4
	427	430	440	464			22 C 12	

HUMAN ARMP FUNCTION DOMAINS

Domai	ns (Am	ino Acid F	Residue)			Functional Characteristic
82	-	100	AA			Hydrophobic
132	-	154	$\mathbf{A}\mathbf{A}$			Hydrophobic
164	-	183	AA			Hydrophobic
195	-	213	AA			Hydrophobic
221	-	238	AA			Hydrophobic
244	-	256	AA			Hydrophobic
281	-	299	AA			Hydrophobic
404	-	428	AA			Hydrophobic
431	_	449	AA			Hydrophobic
115	_	119	AA	(YTPF)	SEQ ID NO: 161	Phosphorylation Site
353	-	356	AA	(STPC)	SEQ ID NO: 162	Phosphorylation Site
300	-	385	AA	-		Acid Rich Domain
						Possible Metal Binding
						Domain

ANTIGENIC SITES INCLUDING AMINO ACID RESIDUES

27	-	44
46	-	48
50	-	60
66	-	67
107	-	111
120	-	121
125	•	126
155	-	160
185	-	189
214	-	223
220		230
240	-	245
267	-	269
273	-	282
300	-	370
400	-	420

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MUTATION	ENZYME (effect of mutation)	AMPLIFICATION 0440 #1	AMPLIFICATION 0440 #2	ALLELE-SPECIFIC 0440
M146LEU	Bsphi (destroy)	910 (170-S182F) TCACAGAAGATACCG AGACT (SEQ ID NO:176)	911 (170-S182) R CCCAACCATAAGAAG AACAG (SEQ ID NO:177)	
MIS 164 Ary	Nia III (destroy)	927 (intronic) TCTGTACTTTTTAAG GGTTGTG (SEQ ID NO:178)	928 ACTICAGAGTAATTC ATCANCA (SEQ ID NO:179)	
Ala 246	Dic I (create)	849 * GACTCCAGCAGGCAT ATCT (SEQ ID NO:80)	892 TGAAATCACAGCCAA GATGAG (SEQ ID NO:130)	
Leu 286 Val.	Pvu II (create)	952 GATGAGACAAGTNCC NTGAA (SEQ ID NO:181) 945 TTAGTGGCTGTTTNG TGTCC (SEQ ID NO:182)	951 CACCCAT'ITACAAGT TTAGC (SEQ ID NO:183)	
Cys 410 Tys	Allele specific ligo	893 GTGTGGCCAGGGTAG AGAACT (SEQ ID NO:128)	885 TGGAGACTGGAACAC AAC (SEQ ID NO:127)	CCATAGCCTGTTTCGTAGC (SEQ ID NO:131) 890 = WT CCATAGCCTATTTCGTAGC (SEQ ID NO:132) 891 = MUT

POSTTICIN OF EXONS AND INTRON-EXON BOUNDARIES OF THE ARYP GENE

CONTRACTOR SEQU	IENC	Corresponding Genom	IC SEQUENCE
ARMP (917 ve)	Transcipt ID CC44 væ	Genomic sequence file D & position of each	Commen
I-113bp	N/A	917-936.gen @ 731-834bp	Alternate 5'UTR
NA	1-4225p	917-936.gen @ 1067-1475bp	Alternate S'UTR
114-195 bp	423-500bp	932-943.zez @ 589-671bp	
19 6- 33 <i>5</i> ap	501-637bp	932-943.zez @ 759-899bp	12bp Variably spliced
337-586 bp	633-883bp	901-912_zes @ 787-1037bp	
587-730bp	\$84-105.5bp	910-915.gen @ 1134-1278bp	MI46L mutation
731-795 cp	1027-109250	925-913.gen @ 413-478bp	H163R mussion
796-1017 bp	1093-1314bp	849-3923== @ 336-558bo	A246E mumica
1018-1116bp	1315-1413bp	951-952-gen @ 312-412hp	L236V mutation, variable spi
1117-1204bp	1414-150120	983-1011.2m @ 61-149bp	
1205-1377bp	1502-1674bp	874-984.gen @ 452-52500	
1378-1497ap	1674-1794bp	885-1012.zm @ 431-550bp	C410Y muzzion
149 3- 2760 00	1795-3060bp	930-919.ze @ -10bp-and	lest AA, STOP, 3"UTR

MUTATIONS AND POLYMORPHISMS IN THE ARMP GENE

Nucleotide # in ARMP.UPD	Amino acid # in ARMP.PRO	Comment
A->C ₆₈₄	Met l 46Leu	Pathogenic, Unique to AD affected.
A->G ₇₃₆	His163Arg	
C->A ₉₈₅	Ala246Glu	n .
C->T ₁₀₂₇	Ala260Val	•
C->T ₁₁₀₃	Ala285Val	•
C->G ₁₁₀₄	Leu286Val	*
C->G ₁₄₂₂	Leu392Vaj	•
G->A ₁₄₇₇	Cys410Tyr	•
G->T ₁₆₃	Phe205Leu	Polymorphism in normals
C->A ₁₇₀₀	non-coding	3'UTR polymorphism
G->A ₂₄₀₁	non-coding	•
delC ₂₄₂₉	non-coding	•

25 - 1	1 METPHANDSEREVCCERTSENSARESPTYREC-QEGROOPEDGEHTAGWANGERGED	\$5			
5182	-	31			
25 -1	S6 G-2EDFDRTVCS-GV9GL99CL	92			
8182		16			
	;				
25 • 1	93 VPVTLCMTVVVATIXEVRPTTEIRCOLTTTPFTEUTPEVQQULLEEVLETLINGSVI	149			
\$182	67 YEVTLOWYVATIXEVEFTTROCGLITTFFTEUTETVOGRALHSTLHAADUSVI	143			
25 -1	150 VYHTIFLVVLYXTRCTXFIBORLINSSLIGLFLFTTTYLGEVLKTYNVANDYFTL-L	205			
5182		200			
-	T t				
	Y I				
25 - 1	206 LTVWNPQAVGRVCIRWEGPLVLOGAYLINE SALMALVPIRFLPRWARWYLGA-18V	261			
5182	201 LT. MINECYVICH LA TEMPERATURE COLUMN TO THE SALVANIVE STREET AND THE AVISY	235			
25 - 1	262 YDLVAVLCPKOPLDGLVETAGEDGRYFPALTYSEAMVVTVGGAELDP\$	310			
8182		312			
R\$ - 1	311 SQGALQLFTDF	345			
\$182]	365			
25-1	346 GTPGENLEREEDGVILG.GDFTFTFVLVGRAAATGSGDGGTTLACFVANLIGLC	400			
6162	166 SSILAGEDPEERGVELGLGDFIFYSVLVGXASATASGDWWTTIACFVAILIGIC	420			
	Ÿ Ÿ				
E5-1	401 LTLLLLAVFIRALPALFISITICLIFFFSTDELVEFFROTLASEQLYI*	448	SEQ	ID	NO:138
518 2			-		NO:2
		-	-		

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